## (19) World Intellectual Property Organization International Bureau



# 

## (43) International Publication Date 20 June 2002 (20.06.2002)

#### **PCT**

## (10) International Publication Number WO 02/48458 A1

(51) International Patent Classification<sup>7</sup>: 17/71

D21H 17/45,

- (21) International Application Number: PCT/US01/48598
- (22) International Filing Date: 7 December 2001 (07.12.2001)
- (25) Filing Language:

English

(26) Publication Language:

English

- (30) Priority Data: 60/256,002 15 December 2000 (15.12.2000) U
- (71) Applicant: THE PROCTER & GAMBLE COMPANY [US/US]; One Procter & Gamble Plaza, Cincinnati, OH 45202 (US).
- (72) Inventors: BARNHOLTZ, Steven, Lee; 5835 Olympia Fields Court, West Chester, OH 45069 (US). VINSON, Kenneth, Douglas; 303 Wyoming Avenue, Cincinnati, OH 45215 (US). COFFARO, Paul, Joseph; 4459 Hickory Bark Court, Cincinnati, OH 45247 (US). MACKEY, Larry, Neil; 5856 Crestview Avenue, Fairfield, OH 45014 (US). HAMILTON, Amy, Jo; 5646 Sante Fe Trail, Mason, OH 45040 (US). WAHL, Errol, Hoffman; 8021 Deershadow Lane, Cincinnati, OH 45242 (US). FRANKENBACH, Gayle, Marie; 10010 Voyager Lane, Cincinnati, OH 45252 (US). WU, Yenchun; 59 Woodstock Drive, Fairfield, OH 45014 (US).

- (74) Agents: REED, T., David et al.; The Procter & Gamble Company, 5299 Spring Grove Avenue, Cincinnati, OH 45217-1087 (US).
- (81) Designated States (national): AE, AG, AL, AM, AT, AT (utility model), AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, CZ (utility model), DE, DE (utility model), DK, DK (utility model), DM, DZ, EC, EE, EE (utility model), ES, FI, FI (utility model), GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PH, PL, PT, RO, RU, SD, SE, SG, SI, SK, SK (utility model), SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW.
- (84) Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

#### Published:

- with international search report
- before the expiration of the time limit for amending the claims and to be republished in the event of receipt of amendments

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

2/48458

(54) Title: SOFT TISSUE PAPER HAVING A SOFTENING COMPOSITION CONTAINING AN EXTENSIONAL VISCOSITY MODIFIER DEPOSITED THEREON

(57) Abstract: Disclosed is a composition for softening an absorbent tissue and tissue structures softened using the composition. The composition includes an effective amount of a softening active ingredient; a vehicle in which the softening active ingredient is dispersed; an electrolyte dissolved in the vehicle; a bilayer disrupter and a high polymer. The electrolyte and the bilayer disrupter cooperate to cause the viscosity of the composition to be less than the viscosity of a dispersion of the softening active ingredient in the vehicle alone. The high polymer adds "stringiness" to the composition opening the air pressure operating window for spray application of the softening composition. Preferably, the softening active ingredient is a quaternary ammonium compound with the formula:  $(R_1)_{4-m} - N^+ - [(CH_2)_n - Y - R_{3lm} X]$  the vehicle is water, the electrolyte is calcium chloride, the bilayer disrupter is a nonionic surfactant, and the high molecular weight polymer is a nonionic polyacrylamide.

## SOFT TISSUE PAPER HAVING A SOFTENING COMPOSITION CONTAINING AN EXTENSIONAL VISCOSITY MODIFIER DEPOSITED THEREON

5

10

#### CROSS REFERENCE

This application claims the benefit of U.S. Provisional Application No. 60/256,002 filed December 15, 2000 and claims priority to U.S. patent application Serial No. 09/413,578, filed in the names of Vinson, et al. on October 6, 1999.

15

#### **TECHNICAL FIELD**

This invention relates, in general, to softening tissue paper; and more specifically, to a composition which may be applied to the surface of tissue paper for enhancing the softness thereof.

#### BACKGROUND OF THE INVENTION

20

25

30

Sanitary paper tissue products are widely used. Such items are commercially offered in formats tailored for a variety of uses such as facial tissues, toilet tissues and absorbent towels.

All of these sanitary products share a common need, specifically to be soft to the touch. Softness is a complex tactile impression evoked by a product when it is stroked against the skin. The purpose of being soft is so that these products can be used to cleanse the skin without being irritating. Effectively cleansing the skin is a persistent personal hygiene problem for many people. Objectionable discharges of urine, menses, and fecal matter from the perineal area or otorhinolaryngogical mucus discharges do not always occur at a time convenient for one to perform a thorough cleansing, as with soap and copious amounts of water for example. As a substitute for thorough cleansing, a wide variety of tissue and toweling products are offered to aid in the task of removing from the skin and retaining such discharges for disposal in a sanitary fashion. Not surprisingly, the use of these products does not approach the level of cleanliness that

can be achieved by the more thorough cleansing methods, and producers of tissue and toweling products are constantly striving to make their products compete more favorably with thorough cleansing methods.

Shortcomings in tissue products for example cause many to stop cleaning before the skin is completely cleansed. Such behavior is often prompted by the harshness of the tissue, as continued rubbing with a harsh implement can abrade the sensitive skin and cause severe pain. The alternative, leaving the skin partially cleansed, is chosen even though this often causes malodors to emanate and can cause staining of undergarments, and over time can cause skin irritations as well.

Disorders of the anus, for example hemorrhoids, render the perianal area extremely sensitive and cause those who suffer such disorders to be particularly frustrated by the need to clean their anus without prompting irritation.

Another notable case which prompts frustration is the repeated nose blowing necessary when one has a cold. Repeated cycles of blowing and wiping can culminate in a sore nose even when the softest tissues available today are employed.

Accordingly, making soft tissue and toweling products which promote comfortable cleaning without performance impairing sacrifices has long been the goal of the engineers and scientists who are devoted to research into improving tissue paper. There have been numerous attempts to reduce the abrasive effect, i.e., improve the softness of tissue products.

One area that has been exploited in this regard has been to select and modify cellulose fiber morphologies and engineer paper structures to take optimum advantages of the various available morphologies. Applicable art in this area includes: Vinson et. al. in U.S. Patent 5,228,954, issued July 20, 1993, Vinson in U.S. Patent 5,405,499, issued April 11, 1995, Cochrane et al. in U.S. Patent 4,874,465 issued October 17, 1989, and Hermans, et. al. in U.S. Statutory Invention Registration H1672, published on August 5, 1997, all of which disclose methods for selecting or upgrading fiber sources to tissue and toweling of superior properties. Applicable art is further illustrated by Carstens in U.S. Patent 4,300,981, issued November 17, 1981, which discusses how fibers can be incorporated to be compliant to paper structures so that they have maximum softness potential. While such techniques as illustrated by these prior art examples are recognized broadly, they can only offer some limited potential to make tissues truly effective comfortable cleaning implements.

5

10

15

20

25

Another area which has received a considerable amount of attention is the addition of chemical softening agents (also referred to herein as "chemical softeners") to tissue and toweling products.

As used herein, the term "chemical softening agent" refers to any chemical ingredient which improves the tactile sensation perceived by the consumer who holds a particular paper product and rubs it across the skin. Although somewhat desirable for towel products, softness is a particularly important property for facial and toilet tissues. Such tactilely perceivable softness can be characterized by, but is not limited to, friction, flexibility, and smoothness, as well as subjective descriptors, such as a feeling like lubricious, velvet, silk or flannel. Suitable materials include those which impart a lubricious feel to tissue. This includes, for exemplary purposes only, basic waxes such as paraffin and beeswax and oils such as mineral oil and silicone oil as well as petrolatum and more complex lubricants and emollients such as quaternary ammonium compounds with long alkyl chains, functional silicones, fatty acids, fatty alcohols and fatty esters.

The field of work in the prior art pertaining to chemical softeners has taken two paths. The first path is characterized by the addition of softeners to the tissue paper web during its formation either by adding an attractive ingredient to the vats of pulp which will ultimately be formed into a tissue paper web, to the pulp slurry as it approaches a paper making machine, or to the wet web as it resides on a Fourdrinier cloth or dryer cloth on a paper making machine.

The second path is categorized by the addition of chemical softeners to tissue paper web after the web is dried. Applicable processes can be incorporated into the paper making operation as, for example, by spraying onto the dry web before it is wound into a roll of paper.

Exemplary art related to the former path categorized by adding chemical softeners to the tissue paper prior to its assembly into a web includes U S. Patent 5,264,082, issued to Phan and Trokhan on November 23, 1993, incorporated herein by reference. Such methods have found broad use in the industry especially when it is desired to reduce the strength which would otherwise be present in the paper and when the papermaking process, particularly the creping operation, is robust enough to tolerate incorporation of the bond inhibiting agents. However, there are problems associated with these methods, well known to those skilled in the art. First, the location of the chemical softener is not controlled; it is spread as broadly through the paper structure as the fiber furnish to which it is applied. In addition, there is a loss of paper strength accompanying use of these additives. While not being bound by theory, it is widely believed that the additives tend to inhibit the formation of fiber to fiber hydrogen bonds. There also can be a loss of control of the sheet as it is creped from the Yankee dryer. Again, a widely believed theory

5

10

15

20

25

is that the additives interfere with the coating on the Yankee dryer so that the bond between the wet web and the dryer is weakened. Prior art such as U.S. Patent 5,487,813, issued to Vinson, et. al., January 30, 1996, incorporated herein by reference, discloses a chemical combination to mitigate the before mentioned effects on strength and adhesion to the creping cylinder; however, there still remains a need to incorporate a chemical softener into a paper web in a targeted fashion with minimal effect on web strength and interference with the production process.

Further exemplary art related to the addition of chemical softeners to the tissue paper web during its formation includes U.S. Patent 5,059,282, issued to Ampulski, et. al. on October 22, 1991 incorporated herein by reference. The Ampulski patent discloses a process for adding a polysiloxane compound to a wet tissue web (preferably at a fiber consistency between about 20% and about 35%). Such a method represents an advance in some respects over the addition of chemicals into the slurry vats supplying the papermaking machine. For example, such means target the application to one of the web surfaces as opposed to distributing the additive onto all of the fibers of the furnish. However, such methods fail to overcome the primary disadvantages of the addition of chemical softeners to the wet end of the papermaking machine, namely the strength effects and the effects on the coating of the Yankee dryer, should such a dryer be employed.

Because of the aforementioned effects on strength and disruption of the papermaking process, considerable art has been devised to apply chemical softeners to already-dried paper webs either at the so-called dry end of the papermaking machine or in a separate converting operation subsequent to the papermaking step. Exemplary art from this field includes U. S. Patent 5,215,626, issued to Ampulski, et. al. on June 1, 1993; U. S. Patent 5,246,545, issued to Ampulski, et. al. on September 21, 1993; U. S. Patent 5,525,345, issued to Warner, et. al. on June 11, 1996, and U.S. Patent application Serial No. 09/053,319 filed in the name of Vinson, et al. on April 1, 1998 all incorporated herein by reference. The 5,215,626 Patent discloses a method for preparing soft tissue paper by applying a polysiloxane to a dry web. The 5,246,545 Patent discloses a similar method utilizing a heated transfer surface. The Warner Patent discloses methods of application including roll coating and extrusion for applying particular compositions to the surface of a dry tissue web. Finally, the Vinson, et al. application discloses compositions that are particularly suitable for surface application onto a tissue web.

While each of these references represent advances over the previous so-called wet end methods, particularly with regard to eliminating the degrading effects on the papermaking process, there remains a need for providing a softening composition that has minimal effect on the

5

10

15

20

25

strength properties of a tissue web. One of the most important physical properties related to softness is generally considered by those skilled in the art to be the strength of the web. Application of a softening composition generally causes a reduction in strength of a tissue web (Strength is the ability of the product, and its constituent webs, to maintain physical integrity and to resist tearing, bursting, and shredding under use conditions). This reduction is believed to result from a disruption of hydrogen bonds between the papermaking fibers that are formed as a result of the papermaking process. Achieving high softness without degrading strength has long been recognized as a means of providing improved tissue products.

Surface application of softening compositions has been found to be particularly useful in providing such softness improvements without substantially degrading the strength of the soft tissue paper product. For example, the parent of the present application discloses compositions that are particularly useful in providing such strong, softened tissue products. Thus, there is a continuing need for commercialization of such soft, strong tissue having a surface applied softening composition.

Commercial production of such strong, softened tissue offers particular challenges. For example, process operation and hygiene can be severely affected if the softening composition is not applied using a suitable process. As noted above with respect to US Patent 5,245,545, the art has applied softening compositions to certain of the rolls in the dry end of a paper machine for transfer to a web surface. While such roll application can successfully apply a softening composition to dried tissue, there are certain process risks. One such risk is roll wrapping in the event of a break in the web downstream of the applicator roll. When such a break occurs, there may be a substantial reduction in takeaway tension on the web at the applicator roll. On loss of takeaway tension, the softening composition can cause the web to adhere to the applicator roll because the takeaway tension is no longer available to overcome adhesive forces due to the softening composition. If such adhesion occurs, there is a high likelihood that the roll will wrap.

Spray application of the softening composition can overcome process operation issues, such as roll wrapping, because the softening composition can be applied to the web in a manner so there is little or no contact between the composition and process rolls before the composition has had a chance to set up. Spray application has its own set of issues, however. One such issue is that, while a minimum air pressure is required to create an spray pattern, there is also a maximum air pressure because particle aerosolization (i.e. creation of very small particles by the spray apparatus) increases with air pressure. The maximum air pressure is defined by the pressure where

5

10

15

20

25

PCT/US01/48598

WO 02/48458

10

15

20

25

such aerosolization results in unacceptable line hygiene (i.e. too much of the softening composition is carried away from the web and deposits on process apparatus).

Accordingly, there is a continuing need for soft tissue paper products having good strength properties. There is also a need for improved softening compositions that can be applied to such tissue products to provide the requisite softness without unacceptably degrading the strength of the product or other important properties thereof. There is a further need for commercial processes capable of producing such products that have acceptable process operation and hygiene.

Such improved products, compositions, and processes are provided by the present invention as is shown in the following disclosure.

#### SUMMARY OF THE INVENTION

The present invention describes softening compositions that, when applied to tissue webs, preferably dried tissue webs, provide soft, strong, absorbent, and aesthetically pleasing tissue paper. The composition is a dispersion comprising:

an effective amount of a softening active ingredient;

a vehicle in which the softening active ingredient is dispersed;

an electrolyte dissolved in the vehicle, the electrolyte causing the viscosity of the composition to be less than the viscosity of a dispersion of the softening composition in the vehicle alone;

a bilayer disrupter to further reduce the viscosity of the softening composition; and

a low level of a high polymer that operates so as to increase the uniaxial and biaxial extensional viscosity of the composition without substantially affecting the shear viscosity thereof.

Reduced aerosolization through the use of this compound meaningfully widens the air pressure operating window for spray application thereof.

The amount of softening active applied to the tissue paper is preferably, between about 0.1% and about 10% based on the total weight of the softening composition compared to the total weight of the resulting tissue paper. The resulting tissue paper preferably has a basis weight of from about 10 to about 80 g/m<sup>2</sup> and a fiber density of less than about 0.6 g/cc.

The term "vehicle" as used herein means a fluid that completely dissolves a chemical papermaking additive, or a fluid that is used to emulsify a chemical papermaking additive, or a fluid that is used to suspend a chemical papermaking additive. The vehicle may also serve as a

carrier that contains a chemical additive or aids in the delivery of a chemical papermaking additive. All references are meant to be interchangeable and not limiting. The dispersion is the fluid containing the chemical papermaking additive. The term "dispersion" as used herein includes true solutions, suspensions, and emulsions. For purposes for this invention, all terms are interchangeable and not limiting. If the vehicle is water or an aqueous solution, then, preferably, the hot web is dried to a moisture level below its equilibrium moisture content (at standard conditions) before being contacted with the composition. However, this process is also applicable to tissue paper at or near its equilibrium moisture content as well.

All percentages, ratios and proportions herein are by weight, unless otherwise specified.

BRIEF DESCRIPTION OF THE FIGURE

While the specification concludes with claims particularly pointing out and distinctly claiming the present invention, it is believed that the present invention will be better understood from the following description in conjunction with the appended example and with the following drawing, in which like reference numbers identify identical elements and wherein:

The figure is a schematic representation illustrating a preferred embodiment of the process of the present invention of adding a softening composition compounds to a tissue web.

The present invention is described in more detail below.

## DETAILED DESCRIPTION OF THE INVENTION

Briefly, the present invention provides a composition which may be applied to a dry tissue web or to a semi-dry tissue web. The resulting tissue paper has enhanced tactilely perceivable softness. The term "dry tissue web" as used herein includes both webs which are dried to a moisture content less than the equilibrium moisture content thereof (overdried-see below) and webs which are at a moisture content in equilibrium with atmospheric moisture. A semi-dry tissue paper web includes a tissue web with a moisture content exceeding its equilibrium moisture content. Most preferably the composition herein is applied to a dry tissue paper web.

The softening composition as well as a method for producing the combination and a method of applying it to tissue are also described.

Surprisingly, it has been found that very low levels of softener additives, e.g. cationic softeners, provide a significant tissue softening effect when applied to the surface of tissue webs in accordance with the present invention. Importantly, it has been found that the levels of softener additives used to soften the tissue paper are low enough that the tissue paper retains high wettability. Furthermore, because the softening composition has a high active level when the

5

10

15

20

25

softening composition is applied, the composition can be applied to dry tissue webs without requiring further drying of the tissue web. Further, since the softening composition of the present invention contains a minimal level of non-functional ingredients, the composition has a minimal effect on the strength of a tissue web after it has been applied.

As used herein, the term "hot tissue web" refers to a tissue web which is at an elevated temperature relative to room temperature. Preferably the elevated temperature of the web is at least about 43°C., and more preferably at least about 65°C.

The moisture content of a tissue web is related to the temperature of the web and the relative humidity of the environment in which the web is placed. As used herein, the term "overdried tissue web" refers to a tissue web that is dried to a moisture content less than its equilibrium moisture content at standard test conditions of 23°C and 50% relative humidity. The equilibrium moisture content of a tissue web placed in standard testing conditions of 23°C and 50% relative humidity is approximately 7%. A tissue web of the present invention can be overdried by raising it to an elevated temperature through use of drying means known to the art such as a Yankee dryer or through air drying. Preferably, an overdried tissue web will have a moisture content of less than 7%, more preferably from about 0 to about 6%, and most preferably, a moisture content of from about 0 to about 3%, by weight.

Paper exposed to the normal environment typically has an equilibrium moisture content in the range of 5 to 8%. When paper is dried and creped the moisture content in the sheet is generally less than 3%. After manufacturing, the paper absorbs water from the atmosphere. In the preferred process of the present invention, advantage is taken of the low moisture content in the paper as it leaves the doctor blade as it is removed from the Yankee dryer (or the low moisture content of similar webs as such webs are removed from alternate drying means if the process does not involve a Yankee dryer).

In a preferred embodiment, the composition of the present invention is applied to an overdried tissue web shortly after it is separated from a drying means and before it is wound onto a parent roll. Alternatively, the composition of the present invention may be applied to a semi-dry tissue web, for example while the web is on the Fourdrinier cloth, on a drying felt or fabric, or while the web is in contact with the Yankee dryer or other alternative drying means. Finally, the composition can also be applied to a dry tissue web in moisture equilibrium with its environment as the web is unwound from a parent roll as for example during an off-line converting operation.

#### Tissue Paper

5

10

15

20

25

The present invention is applicable to tissue paper in general, including but not limited to: conventionally felt-pressed tissue paper; pattern densified tissue paper such as exemplified by Sanford-Sisson and its progeny; and high-bulk, uncompacted tissue paper such as exemplified by Salvucci. The tissue paper may be of a homogenous or multilayered construction; and tissue paper products made therefrom may be of a single-ply or multi-ply construction. The tissue paper preferably has a basis weight of between about 10 g/m² and about 80 g/m², and density of about 0.60 g/cc or less. Preferably, the basis weight will be below about 35 g/m² or less; and the density will be about 0.30 g/cc or less. Most preferably, the density will be between about 0.04 g/cc and about 0.20 g/cc.

Conventionally pressed tissue paper and methods for making such paper are known in the art. Such paper is typically made by depositing a papermaking furnish on a foraminous forming wire. This forming wire is often referred to in the art as a Fourdrinier wire. Once the furnish is deposited on the forming wire, it is referred to as a web. Overall, water is removed from the web by vacuum, mechanical pressing and thermal means. The web is dewatered by pressing the web and by drying at elevated temperature. The particular techniques and typical equipment for making webs according to the process just described are well known to those skilled in the art. In a typical process, a low consistency pulp furnish is provided in a pressurized headbox. The headbox has an opening for delivering a thin deposit of pulp furnish onto the Fourdrinier wire to form a wet web. The web is then typically dewatered to a fiber consistency of between about 7% and about 45% (total web weight basis) by vacuum dewatering and further dried by pressing operations wherein the web is subjected to pressure developed by opposing mechanical members, for example, cylindrical rolls. The dewatered web is then further pressed and dried by a stream drum apparatus known in the art as a Yankee dryer. Pressure can be developed at the Yankee dryer by mechanical means such as an opposing cylindrical drum pressing against the web. Multiple Yankee dryer drums may be employed, whereby additional pressing is optionally incurred between the drums. The tissue paper structures which are formed are referred to hereinafter as conventional, pressed, tissue paper structures. Such sheets are considered to be compacted, since the web is subjected to substantial overall mechanical compression forces while the fibers are moist and are then dried while in a compressed state. The resulting structure is strong and generally of singular density, but very low in bulk, absorbency and in softness.

Pattern densified tissue paper is characterized by having a relatively high-bulk field of relatively low fiber density and an array of densified zones of relatively high fiber density. The high-bulk field is alternatively characterized as a field of pillow regions. The densified zones are alternatively referred to as knuckle regions. The densified zones may be discretely spaced within

5

10

15

20

25

the high-bulk field or may be interconnected, either fully or partially, within the high-bulk field. Preferred processes for making pattern densified tissue webs are disclosed in U.S. Patent 3,301,746, issued to Sanford and Sisson on January 31, 1967, U.S. Patent 3,974,025, issued to Ayers on August 10, 1976, and U.S. Patent 4,191,609, issued to on March 4, 1980, and U.S. Patent 4,637,859, issued to on January 20, 1987; the disclosure of each of which is incorporated herein by reference.

5

10

15

20

25

30

In general, pattern densified webs are preferably prepared by depositing a papermaking furnish on a foraminous forming wire such as a Fourdrinier wire to form a wet web and then juxtaposing the web against an array of supports as it is transferred from the forming wire to a structure comprising such supports for further drying. The web is pressed against the array of supports, thereby resulting in densified zones in the web at the locations geographically corresponding to the points of contact between the array of supports and the wet web. The remainder of the web not compressed during this operation is referred to as the high-bulk field. This high-bulk field can be further dedensified by application of fluid pressure, such as with a vacuum type device or a blow-through dryer, or by mechanically pressing the web against the array of supports. The web is dewatered, and optionally predried, in such a manner so as to substantially avoid compression of the high-bulk field. This is preferably accomplished by fluid pressure, such as with a vacuum type device or blow-through dryer, or alternately by mechanically pressing the web against an array of supports wherein the high-bulk field is not compressed. The operations of dewatering, optional predrying and formation of the densified zones may be integrated or partially integrated to reduce the total number of processing steps performed. Subsequent to formation of the densified zones, dewatering, and optional predrying, the web is dried to completion, preferably still avoiding mechanical pressing. Preferably, from about 8% to about 65% of the tissue paper surface comprises densified knuckles, the knuckles preferably having a relative density of at least 125% of the density of the high-bulk field.

The structure comprising an array of supports is preferably an imprinting carrier fabric having a patterned displacement of knuckles which operate as the array of supports which facilitate the formation of the densified zones upon application of pressure. The pattern of knuckles constitutes the array of supports previously referred to. Imprinting carrier fabrics are disclosed in U.S. Patent 3,301,746, issued to Sanford and Sisson on January 31, 1967, U.S. Patent 3,821,068, issued to Salvucci, Jr. et al. on May 21, 1974, U.S. Patent 3,974,025, issued to Ayers on August 10, 1976, U.S. Patent 3,573,164, issued to Friedberg, et al. on March 30, 1971, U.S. Patent 3,473,576, issued to Amneus on October 21, 1969, U.S. Patent 4,239,065, issued to

Trokhan on December 16, 1980, and U.S. Patent 4,528,239, issued to Trokhan on July 9, 1985, the disclosure of each of which is incorporated herein by reference.

Preferably, the furnish is first formed into a wet web on a foraminous forming carrier, such as a Fourdrinier wire. The web is dewatered and transferred to an imprinting fabric. The furnish may alternately be initially deposited on a foraminous supporting carrier which also operates as an imprinting fabric. Once formed, the wet web is dewatered and, preferably, thermally predried to a selected fiber consistency of between about 40% and about 80%. Dewatering is preferably performed with suction boxes or other vacuum devices or with blow-through dryers. The knuckle imprint of the imprinting fabric is impressed in the web as discussed above, prior to drying the web to completion. One method for accomplishing this is through application of mechanical pressure. This can be done, for example, by pressing a nip roll which supports the imprinting fabric against the face of a drying drum, such as a Yankee dryer, wherein the web is disposed between the nip roll and drying drum. Also, preferably, the web is molded against the imprinting fabric prior to completion of drying by application of fluid pressure with a vacuum device such as a suction box, or with a blow-through dryer. Fluid pressure may be applied to induce impression of densified zones during initial dewatering, in a separate, subsequent process stage, or a combination thereof.

Uncompacted, non pattern-densified tissue paper structures are described in U.S. Patent 3,812,000 issued to Joseph L. Salvucci, Jr. and Peter N. Yiannos on May 21, 1974, and U.S. Patent 4,208,459, issued to Henry E. Becker, Albert L. McConnell, and Richard Schutte on Jun. 17, 1980, both of which are incorporated herein by reference. In general, uncompacted, non pattern-densified tissue paper structures are prepared by depositing a papermaking furnish on a foraminous forming wire such as a Fourdrinier wire to form a wet web, draining the web and removing additional water without mechanical compression until the web has a fiber consistency of at least 80%, and creping the web. Water is removed from the web by vacuum dewatering and thermal drying. The resulting structure is a soft but weak high-bulk sheet of relatively uncompacted fibers. Bonding material is preferably applied to portions of the web prior to creping.

The softening composition of the present invention can also be applied to uncreped tissue paper. Uncreped tissue paper, a term as used herein, refers to tissue paper which is non-compressively dried, most preferably by through air drying. Resultant through air dried webs are pattern densified such that zones of relatively high density are dispersed within a high bulk field,

5

10

15

20

25

including pattern densified tissue wherein zones of relatively high density are continuous and the high bulk field is discrete.

To produce uncreped tissue paper webs, an embryonic web is transferred from the foraminous forming carrier upon which it is laid, to a slower moving, high fiber support transfer fabric carrier. The web is then transferred to a drying fabric upon which it is dried to a final dryness. Such webs can offer some advantages in surface smoothness compared to creped paper webs.

The techniques to produce uncreped tissue in this manner are taught in the prior art. For example, Wendt, et. al. in European Patent Application 0 677 612A2, published October 18, 1995 and incorporated herein by reference, teach a method of making soft tissue products without creping. In another case, Hyland, et. al. in European Patent Application 0 617 164 A1, published September 28, 1994 and incorporated herein by reference, teach a method of making smooth uncreped through air dried sheets. Finally, Farrington, et. al. in U.S. Patent 5,656,132 published August 12, 1997, the disclosure of which is incorporated herein by reference, describes the use of a machine to make soft through air dried tissues without the use of a Yankee.

#### **Furnish**

5

10

15

20

25

30

#### Papermaking Fibers

The papermaking fibers utilized for the present invention will normally include fibers derived from wood pulp. Other cellulosic fibrous pulp fibers, such as cotton linters, bagasse, etc., can be utilized and are intended to be within the scope of this invention. Synthetic fibers, such as rayon, polyethylene and polypropylene fibers, may also be utilized in combination with natural cellulosic fibers. One exemplary polyethylene fiber which may be utilized is Pulpex®, available from Hercules, Inc. (Wilmington, DE).

Applicable wood pulps include chemical pulps, such as Kraft, sulfite, and sulfate pulps, as well as mechanical pulps including, for example, groundwood, thermomechanical pulp and chemically modified thermomechanical pulp. Chemical pulps, however, are preferred since they impart a superior tactile sense of softness to tissue sheets made therefrom. Pulps derived from both deciduous trees (hereinafter, also referred to as "hardwood") and coniferous trees (hereinafter, also referred to as "softwood") may be utilized. Also applicable to the present invention are fibers derived from recycled paper, which may contain any or all of the above categories as well as other non-fibrous materials such as fillers and adhesives used to facilitate the original papermaking.

## Optional Chemical Additives

5

10

15

20

25

30

Other materials can be added to the aqueous papermaking furnish or the embryonic web to impart other desirable characteristics to the product or improve the papermaking process so long as they are compatible with the chemistry of the softening composition and do not significantly and adversely affect the softness or strength character of the present invention. The following materials are expressly included, but their inclusion is not offered to be all-inclusive. Other materials can be included as well so long as they do not interfere or counteract the advantages of the present invention.

It is common to add a cationic charge biasing species to the papermaking process to control the zeta potential of the aqueous papermaking furnish as it is delivered to the papermaking process. These materials are used because most of the solids in nature have negative surface charges, including the surfaces of cellulosic fibers and fines and most inorganic fillers. One traditionally used cationic charge biasing species is alum. More recently in the art, charge biasing is done by use of relatively low molecular weight cationic synthetic polymers preferably having a molecular weight of no more than about 500,000 and more preferably no more than about 200,000, or even about 100,000. The charge densities of such low molecular weight cationic synthetic polymers are relatively high. These charge densities range from about 4 to about 8 equivalents of cationic nitrogen per kilogram of polymer. An exemplary material is Cypro 514°, a product of Cytec, Inc. of Stamford, CT. The use of such materials is expressly allowed within the practice of the present invention.

The use of high surface area, high anionic charge microparticles for the purposes of improving formation, drainage, strength, and retention is taught in the art. See, for example, U. S. Patent, 5,221,435, issued to Smith on June 22, 1993, the disclosure of which is incorporated herein by reference. Common materials for this purpose are silica colloid, or bentonite clay. The incorporation of such materials is expressly included within the scope of the present invention.

If permanent wet strength is desired, the group of chemicals: including polyamide-epichlorohydrin, polyacrylamides, styrene-butadiene lattices; insolubilized polyvinyl alcohol; urea-formaldehyde; polyethyleneimine; chitosan polymers and mixtures thereof can be added to the papermaking furnish or to the embryonic web. Preferred resins are cationic wet strength resins, such as polyamide-epichlorohydrin resins. Suitable types of such resins are described in U.S. Patents 3,700,623, issued on October 24, 1972, and 3,772,076, issued on November 13, 1973, both to Keim, the disclosure of both being hereby incorporated by reference. One

commercial source of useful polyamide-epichlorohydrin resins is Hercules, Inc. of Wilmington, Delaware, which markets such resin under the mark Kymene 557H<sup>®</sup>.

Many paper products must have limited strength when wet because of the need to dispose of them through toilets into septic or sewer systems. If wet strength is imparted to these products, fugitive wet strength, characterized by a decay of part or all of the initial strength upon standing in presence of water, is preferred. If fugitive wet strength is desired, the binder materials can be chosen from the group consisting of dialdehyde starch or other resins with aldehyde functionality such as Co-Bond 1000® offered by National Starch and Chemical Company of Scarborough, ME; Parez 750® offered by Cytec of Stamford, CT; and the resin described in U.S. Patent 4,981,557, issued on January 1, 1991, to Bjorkquist, the disclosure of which is incorporated herein by reference, and other such resins having the decay properties described above as may be known to the art.

If enhanced absorbency is needed, surfactants may be used to treat the tissue paper webs of the present invention. The level of surfactant, if used, is preferably from about 0.01% to about 2.0% by weight, based on the dry fiber weight of the tissue web. The surfactants preferably have alkyl chains with eight or more carbon atoms. Exemplary anionic surfactants include linear alkyl sulfonates and alkylbenzene sulfonates. Exemplary nonionic surfactants include alkylglycosides including alkylglycoside esters such as Crodesta SL-40® which is available from Croda, Inc. (New York, NY); alkylglycoside ethers as described in U.S. Patent 4,011,389, issued to Langdon, et al. on March 8, 1977; and alkylpolyethoxylated esters such as Pegosperse 200 ML available from Glyco Chemicals, Inc. (Greenwich, CT) and IGEPAL RC-520® available from Rhone Poulenc Corporation (Cranbury, NJ). Alternatively, cationic softener active ingredients with a high degree of unsaturated (mono and/or poly) and/or branched chain alkyl groups can greatly enhance absorbency.

While the essence of the present invention is the presence of a softening agent composition deposited on the tissue web surface, the invention also expressly includes variations in which chemical softening agents are added as a part of the papermaking process. For example, chemical softening agents may be included by wet end addition. Preferred chemical softening agents comprise quaternary ammonium compounds including, but not limited to, the well-known dialkyldimethylammonium salts (e.g., ditallowdimethylammonium chloride, ditallowdimethylammonium methyl sulfate, di(hydrogenated tallow)dimethyl ammonium chloride, etc.). Particularly preferred variants of these softening agents include mono or diester variations of the before mentioned dialkyldimethylammonium salts and ester quaternaries made

5

10

15

20

25

from the reaction of fatty acid and either methyl diethanol amine and/or triethanol amine, followed by quaternization with methyl chloride or dimethyl sulfate.

Another class of papermaking-added chemical softening agents comprise the well-known organo-reactive polydimethyl siloxane ingredients, including the most preferred amino functional polydimethyl siloxane.

Filler materials may also be incorporated into the tissue papers of the present invention. U.S. Patent 5,611,890, issued to Vinson et al. on March 18, 1997, and, incorporated herein by reference discloses filled tissue paper products that are acceptable as substrates for the present invention.

The above listings of optional chemical additives is intended to be merely exemplary in nature, and are not meant to limit the scope of the invention.

### Softening Composition

5

10

15

20

In general, the softening composition of the present invention comprises a dispersion of a softening active ingredient in a vehicle. When applied to tissue paper as described herein, such compositions are effective in softening the tissue paper. Preferably, the softening composition of the present invention has properties (e.g., ingredients, rheology, pH, etc.) permitting easy application thereof on a commercial scale. For example, while certain volatile organic solvents may readily dissolve high concentrations of effective softening materials, such solvents are not desired because of the increased process safety and environmental burden (VOC) concerns raised by such solvents. The following discusses each of the components of the softening composition of the present invention, the properties of the composition, methods of producing the composition, and methods of applying the composition.

#### Components

## Softening Active Ingredients

25 Quaternary compounds having the formula:

$$(R_1)_{4-m} - N^+ - [R_2]_m X^-$$

wherein:

m is 1 to 3;

each R<sub>1</sub> is a C<sub>1</sub>-C<sub>6</sub> alkyl group, hydroxyalkyl group, hydrocarbyl or substituted hydrocarbyl group, alkoxylated group, benzyl group, or mixtures thereof;

each R<sub>2</sub> is a C<sub>14</sub>-C<sub>22</sub> alkyl group, hydroxyalkyl group, hydrocarbyl or substituted hydrocarbyl group, alkoxylated group, benzyl group, or mixtures thereof; and

X is any softener-compatible anion

are suitable for use in the present invention. Preferably, each R<sub>1</sub> is methyl and X<sup>-</sup> is chloride or methyl sulfate. Preferably, each R<sub>2</sub> is C<sub>16</sub>-C<sub>18</sub> alkyl or alkenyl, most preferably each R<sub>2</sub> is straight-chain C<sub>18</sub> alkyl or alkenyl. Optionally, the R<sub>2</sub> substituent can be derived from vegetable oil sources. Several types of the vegetable oils (e.g., olive, canola, safflower, sunflower, etc.) can used as sources of fatty acids to synthesize the quaternary ammonium compound. Branched chain actives (e.g., made from isostearic acid) are also effective.

Such structures include the well-known dialkyldimethylammonium salts (e.g., ditallowdimethylammonium chloride, ditallowdimethylammonium methyl sulfate, di(hydrogenated tallow)dimethyl ammonium chloride, etc.), in which R<sub>1</sub> are methyl groups, R<sub>2</sub> are tallow groups of varying levels of saturation, and X<sup>-</sup> is chloride or methyl sulfate.

As discussed in Swern, Ed. in Bailey's Industrial Oil and Fat Products, Third Edition, John Wiley and Sons (New York 1964), tallow is a naturally occurring material having a variable composition. Table 6.13 in the above-identified reference edited by Swern indicates that typically 78% or more of the fatty acids of tallow contain 16 or 18 carbon atoms. Typically, half of the fatty acids present in tallow are unsaturated, primarily in the form of oleic acid. Synthetic as well as natural "tallows" fall within the scope of the present invention. It is also known that depending upon the product characteristic requirements, the saturation level of the ditallow can be tailored from non hydrogenated (soft) to touch (partially hydrogenated) or completely hydrogenated (hard). All of above-described saturation levels of are expressly meant to be included within the scope of the present invention.

Particularly preferred variants of these softening active ingredients are what are considered to be mono or diester variations of these quaternary ammonium compounds having the formula:

$$(R_1)_{4-m} - N^+ - [(CH_2)_n - Y - R_3]_m X^-$$

wherein

Y is -O-(O)C-, or -C(O)-O-, or -NH-C(O)-, or -C(O)-NH-;

m is 1 to 3;

30 n is 0 to 4;

10

15

20

each R<sub>1</sub> is a C<sub>1</sub>-C<sub>6</sub> alkyl group, hydroxyalkyl group, hydrocarbyl or substituted hydrocarbyl group, alkoxylated group, benzyl group, or mixtures thereof;

each  $R_3$  is a  $C_{13}$ - $C_{21}$  alkyl group, hydroxyalkyl group, hydrocarbyl or substituted hydrocarbyl group, alkoxylated group, benzyl group, or mixtures thereof; and

X is any softener-compatible anion.

Preferably, Y = -O-(O)C-, or -C(O)-O-; m=2; and n=2. Each  $R_1$  substituent is preferably a  $C_1-C_3$ , alkyl group, with methyl being most preferred. Preferably, each  $R_3$  is  $C_{13}$  - $C_{17}$  alkyl and/or alkenyl, more preferably  $R_3$  is straight chain  $C_{15}$  -  $C_{17}$  alkyl and/or alkenyl,  $C_{15}$ - $C_{17}$  alkyl, most preferably each  $R_3$  is straight-chain  $C_{17}$  alkyl. Optionally, the  $R_3$  substituent can be derived from vegetable oil sources. Several types of the vegetable oils (e.g., olive, canola, safflower, sunflower, etc.) can used as sources of fatty acids to synthesize the quaternary ammonium compound. Preferably, olive oils, canola oils, high oleic safflower, and/or high erucic rapeseed oils are used to synthesize the quaternary ammonium compound.

As mentioned above, X<sup>-</sup> can be any softener-compatible anion, for example, acetate, chloride, bromide, methylsulfate, formate, sulfate, nitrate and the like can also be used in the present invention. Preferably X<sup>-</sup> is chloride or methyl sulfate.

Specific examples of ester-functional quaternary ammonium compounds having the structures named above and suitable for use in the present invention include the well-known diester dialkyl dimethyl ammonium salts such as diester ditallow dimethyl ammonium chloride, monoester ditallow dimethyl ammonium chloride, diester ditallow dimethyl ammonium methyl sulfate, diester di(hydrogenated)tallow dimethyl ammonium methyl sulfate, diester di(hydrogenated)tallow dimethyl ammonium chloride, and mixtures thereof. Diester ditallow dimethyl ammonium chloride are particularly preferred. These particular materials are available commercially from Goldschmidt Chemical Corporation of Dublin, OH under the tradename ADOGEN SDMC. Also preferred for their low corrosivity are the methyl sulfate salts of such quaternary cations. These materials are also available in experimental quantities from Goldschmidt Chemical company.

As mentioned above, typically, half of the fatty acids present in tallow are unsaturated, primarily in the form of oleic acid. Synthetic as well as natural "tallows" fall within the scope of the present invention. It is also known that depending upon the product characteristic requirements, the degree of saturation for such tallows can be tailored from non hydrogenated (soft), to partially hydrogenated (touch), or completely hydrogenated (hard). All of above-

5

10

15

20

25

described saturation levels of are expressly meant to be included within the scope of the present invention.

It will be understood that substituents R<sub>1</sub>, R<sub>2</sub> and R<sub>3</sub> may optionally be substituted with various groups such as alkoxyl, hydroxyl, or can be branched. As mentioned above, preferably each R<sub>1</sub> is methyl or hydroxyethyl. Preferably, each R<sub>2</sub> is C<sub>12</sub> - C<sub>18</sub> alkyl and/or alkenyl, most preferably each R<sub>2</sub> is straight-chain C<sub>16</sub> - C<sub>18</sub> alkyl and/or alkenyl, most preferably each R<sub>2</sub> is straight-chain C<sub>18</sub> alkyl or alkenyl. Preferably R<sub>3</sub> is C<sub>13</sub> - C<sub>17</sub> alkyl and/or alkenyl, most preferably R<sub>3</sub> is straight chain C<sub>15</sub> - C<sub>17</sub> alkyl and/or alkenyl. Preferably, X<sup>-</sup> is chloride or methyl sulfate. Furthermore the ester-functional quaternary ammonium compounds can optionally contain up to about 10% of the mono(long chain alkyl) derivatives, e.g.:

5

10

15

20

25

30

as minor ingredients. These minor ingredients can act as emulsifiers and are useful in the present invention.

Other types of suitable quaternary ammonium compounds for use in the present invention are described in U.S. Patent 5,543,067, issued to Phan et al. on August 6, 1996; U.S. Patent 5,538,595, issued to Trokhan et al., on July 23, 1996; U.S. Patent 5,510,000, issued to Phan et al. on April 23, 1996; U.S. Patent 5,415,737, issued to Phan et al., on May 16, 1995; and European Patent Application No. 0 688 901 A2, assigned to Kimberly-Clark Corporation, published December 12, 1995; the disclosure of each of which is incorporated herein by reference.

Di-quat variations of the ester-functional quaternary ammonium compounds can also be used, and are meant to fall within the scope of the present invention. These compounds have the formula:

O 
$$(R_1)_2$$
  $(R_1)_2$  O  $||$   $||$   $||$   $||$   $||$   $||$   $R_1 - C - O - (CH_2)_2 - N^+ - (CH_2)_n - N^+ (CH_2)_2 - O - C - R_3$  2 X

In the structure named above each  $R_1$  is a  $C_1$  -  $C_6$ alkyl or hydroxyalkyl group,  $R_3$  is  $C_{11}$ - $C_{21}$  hydrocarbyl group, n is 2 to 4 and  $X^-$  is a suitable anion, such as an halide (e.g., chloride or bromide) or methyl sulfate. Preferably, each  $R_3$  is  $C_{13}$ - $C_{17}$  alkyl and/or alkenyl, most preferably each  $R_3$  is straight-chain  $C_{15}$  -  $C_{17}$  alkyl and/or alkenyl, and  $R_1$  is a methyl.

Parenthetically, while not wishing to be bound by theory, it is believed that the ester moiety(ies) of the aforementioned quaternary compounds provides a measure of biodegradability to such compounds. Importantly, the ester-functional quaternary ammonium compounds used

herein biodegrade more rapidly than do conventional dialkyl dimethyl ammonium chemical softeners.

The use of quaternary ammonium ingredients as described herein above is most effectively accomplished if the quaternary ammonium ingredient is accompanied by an appropriate plasticizer. The term plasticizer as used herein refers to an ingredient capable of reducing the melting point and viscosity at a given temperature of a quaternary ammonium ingredient. The plasticizer can be added during the quaternizing step in the manufacture of the quaternary ammonium ingredient or it can be added subsequent to the quaternization but prior to the application as a softening active ingredient. The plasticizer is characterized by being substantially inert during the chemical synthesis which acts as a viscosity reducer to aid in the synthesis. Preferred plasticizers are non-volatile polyhydroxy compounds. Preferred polyhydroxy compounds include glycerol and polyethylene glycols having a molecular weight of from about 200 to about 2000, with polyethylene glycol having a molecular weight of from about 200 to about 600 being particularly preferred. When such plasticizers are added during manufacture of the quaternary ammonium ingredient, they comprise between about 5% and about 75% percent of the product of such manufacture. Particularly preferred mixtures comprise between about 15% and about 50% plasticizer.

#### Vehicle

5

10

15

20

25

30

As used herein a "vehicle" is used to dilute the active ingredients of the compositions described herein forming the dispersion of the present invention. A vehicle may dissolve such components (true solution or micellar solution) or such components may be dispersed throughout the vehicle (dispersion or emulsion). The vehicle of a suspension or emulsion is typically the continuous phase thereof. That is, other components of the dispersion or emulsion are dispersed on a molecular level or as discrete particles throughout the vehicle.

For purposes of the present invention, one purpose that the vehicle serves is to dilute the concentration of softening active ingredients so that such ingredients may be efficiently and economically applied to a tissue web. For example, as is discussed below, one way of applying such active ingredients is to spray them onto a roll which then transfers the active ingredients to a moving web of tissue. Typically, only very low levels (e. g. on the order of 2% by weight of the associated tissue) of softening active ingredients are required to effectively improve the tactile sense of softness of a tissue. This means very accurate metering and spraying systems would be required to distribute a "pure" softening active ingredient across the full width of a commercial-scale tissue web.

Another purpose of the vehicle is to deliver the active softening composition in a form in which it is less prone to be mobile with regard to the tissue structure. Specifically, it is desired to apply the composition of the present invention so that the active ingredient of the composition resides primarily on the surface of the absorbent tissue web with minimal absorption into the interior of the web. While not wishing to be bound by theory, the Applicants believe that the interaction of the softening composition with preferred vehicles creates a suspended particle which binds more quickly and permanently than if the active ingredient were to be applied without the vehicle. For example, it is believed that suspensions of quaternary softeners in water assume a liquid crystalline form which can be substantively deposited onto the surface of the fibers of the surface of the tissue paper web. Quaternary softeners applied without the aid of the vehicle, e. g. applied in molten form by contrast tend to wick into the internal of the tissue web.

The Applicants have discovered vehicles and softening compositions comprising such vehicles that are particularly useful for facilitating the application of softening active ingredients to webs of tissue on a commercial scale.

While softening ingredients can be dissolved in a vehicle forming a solution therein, materials that are useful as solvents for suitable softening active ingredients are not commercially desirable for safety and environmental reasons. Therefore, to be suitable for use in the vehicle for purposes of the present invention, a material should be compatible with the softening active ingredients described herein and with the tissue substrate on which the softening compositions of the present invention will be deposited. Further a suitable material should not contain any ingredients that create safety issues (either in the tissue manufacturing process or to users of tissue products using the softening compositions described herein) and not create an unacceptable risk to the environment. Suitable materials for the vehicle of the present invention include hydroxyl functional liquids most preferably water.

#### 25 <u>Electrolyte</u>

5

10

15

20

30

While water is a particularly preferred material for use in the vehicle of the present invention, water alone is not preferred as a vehicle. Specifically, when softening active ingredients of the present invention are dispersed in water at a level suitable for application to a tissue web, the dispersion has an unacceptably high viscosity. While not being bound by theory, the Applicants believe that combining water and the softening active ingredients of the present invention to form such dispersions creates a liquid crystalline phase having a high viscosity. Compositions having such a high viscosity are difficult to apply to tissue webs for softening purposes.

The Applicants have discovered that the viscosity of dispersions of softening active ingredients in water can be substantially reduced, while maintaining a desirable high level of the softening active ingredient in the softening composition by the simple addition of a suitable electrolyte to the vehicle. Again, not being bound by theory, the Applicants believe the electrolyte shields the electrical charge around bilayers and vesicles, reducing interactions, and lowering resistance to movement resulting in a reduction in viscosity of the system. Additionally, again not being bound by theory, the electrolyte can create an osmotic pressure difference across vesicle walls which would tend to draw interior water through the vesicle wall reducing the size of the vesicles and providing more "free" water, again resulting in a decrease in viscosity.

Any electrolyte meeting the general criteria described above for materials suitable for use in the vehicle of the present invention and which is effective in reducing the viscosity of a dispersion of a softening active ingredient in water is suitable for use in the vehicle of the present invention. In particular, any of the known water-soluble electrolytes meeting the above criteria can be included in the vehicle of the softening composition of the present invention. When present, the electrolyte can be used in amounts up to about 25% by weight of the softening composition, but preferably no more than about 15 % by weight of the softening composition. Preferably, the level of electrolyte is between about 0.1% and about 10% by weight of the softening composition based on the anhydrous weight of the electrolyte. Still more preferably, the electrolyte is used at a level of between about 0.3% and about 1.0% by weight of the softening composition. The minimum amount of the electrolyte will be that amount sufficient to provide the desired viscosity. The dispersions typically display a non-Newtonian rheology, and are shear thinning with a desired viscosity generally ranging from about 10 centipoise (cp) up to about 1000 cp, preferably in the range between about 10 and about 200 cp, as measured at 25° C and at a shear rate of 100 sec-1 using the method described in the TEST Methods section below. Suitable electrolytes include the halide, nitrate, nitrite, and sulfate salts of alkali or alkaline earth metals, as well as the corresponding ammonium salts. Other useful electrolytes include the alkali and alkaline earth salts of simple organic acids such as sodium formate and sodium acetate, as well as the corresponding ammonium salts. Preferred inorganic electrolytes include the chloride salts of sodium, calcium, and magnesium. Calcium chloride is a particularly preferred inorganic electrolyte for the softening composition of the present invention. A particularly preferred organic acid salt-based electrolyte is sodium formate.

#### Bilayer Disrupter

5

10

15

20

25

5

10

15

20

25

30

A bilayer disrupter is an essential component of the invention. While, as has been shown above, the vehicle, particularly the electrolyte thereof, performs an essential function in preparing the soft tissue paper webs of the present invention, it is desirable also to limit the amount of vehicle deposited onto a tissue web. As noted above, addition of electrolyte allows an increase in the concentration of softening active ingredient in the softening composition without unduly increasing viscosity. However, if too much electrolyte is used, phase separation can occur. The Applicants have found that adding a bilayer disrupter to the softening composition allows more softening active ingredient to be incorporated therein while maintaining viscosity at an acceptable level. As used herein a "bilayer disrupter" is an organic material that, when mixed with a dispersion of a softening active ingredient in a vehicle, is compatible with at least one of the vehicle or the softening active ingredient and causes a reduction of the viscosity of the dispersion.

Not to be bound by theory, it is believed that bilayer disrupters function by penetrating the pallisade layer of the liquid crystalline structure of the dispersion of the softening active ingredient in the vehicle and disrupting the order of the liquid crystalline structure. Such disruption is believed to reduce the interfacial tension at the hydrophobic-water interface, thus promoting flexibility with a resulting reduction in viscosity. As used herein, the term "pallisade layer", it is meant describe the area between hydrophilic groups and the first few carbon atoms in the hydrophobic layer (M.J Rosen, Surfactants and interfacial phenomena, Second Edition, pages 125 and 126).

In addition to providing the viscosity reduction benefits discussed above, materials suitable for use as a bilayer disrupter should be compatible with other components of the softening composition. For example, a suitable material should not react with other components of the softening composition so as to cause the softening composition to lose softening capability.

Bilayer disrupters useful in the compositions of the present invention are preferably surface active materials. Such materials comprise both hydrophobic and hydrophilic moieties. A preferred hydrophilic moiety is a polyalkoxylated group, preferably a polyethoxylated group. Such preferred bilayer disrupters are used at a level of between about 2% and about 15% of the level of the softening active ingredient. Preferably, the bilayer disrupter is present at a level of between about 3% and about 10% of the level of the softening active ingredient.

Particularly preferred bilayer disrupters are nonionic surfactants derived from saturated and/or unsaturated primary and/or secondary, amine, amide, amine-oxide fatty alcohol, fatty acid, alkyl phenol, and/or alkyl aryl carboxylic acid compounds, each preferably having from about 6 to about 22, more preferably from about 8 to about 18, carbon atoms in a hydrophobic chain,

5

10

15

more preferably an alkyl or alkylene chain, wherein at least one active hydrogen of said compounds is ethoxylated with  $\leq 50$ , preferably  $\leq 30$ , more preferably from about 3 to about 15, and even more preferably from about 5 to about 12, ethylene oxide moieties to provide an HLB of from about 6 to about 20, preferably from about 8 to about 18, and more preferably from about 10 to about 15.

Suitable bilayer disrupters also include nonionic surfactants with bulky head groups selected from:

## a. surfactants having the formula

$$R^{1}$$
-C(O)-Y'-[C( $R^{5}$ )]<sub>m</sub>-CH<sub>2</sub>O( $R_{2}$ O)<sub>z</sub>H

wherein  $R^1$  is selected from the group consisting of saturated or unsaturated, primary, secondary or branched chain alkyl or alkyl-aryl hydrocarbons; said hydrocarbon chain having a length of from about 6 to about 22; Y' is selected from the following groups: -O-; -N(A)-; and mixtures thereof; and A is selected from the following groups: H;  $R^1$ ; -( $R^2$ -O)<sub>z</sub>-H; -( $R^2$ -O)<sub>z</sub>-H; -( $R^2$ -O)<sub>z</sub>-H; phenyl, or substituted aryl, wherein  $0 \le x \le about 3$  and z is from about 5 to about 30; each  $R^2$  is selected from the following groups or combinations of the following groups: -( $R^2$ -O)<sub>z</sub>-H; and m is from about 2 to about 4;

## b. surfactants having the formulas:

$$R^5$$
 $R^5$ 
 $R^5$ 
 $R^5$ 
 $R^5$ 

20

25

wherein Y" = N or O; and each R5 is selected independently from the following:

-H, -OH, -(CH<sub>2</sub>)xCH<sub>3</sub>, -O(OR<sup>2</sup>)<sub>z</sub>-H, -OR<sup>1</sup>, - OC(O)R<sup>1</sup>, and -CH(CH<sub>2</sub>-(OR<sup>2</sup>)<sub>z</sub>-H)-CH<sub>2</sub>-(OR<sup>2</sup>)<sub>z</sub>-C(O) R<sup>1</sup>, x and R<sup>1</sup> are as defined above and  $5 \le z$ , z, and z''  $\le 20$ , more preferably  $5 \le z + z$ , + z,  $\le 20$ , and most preferably, the heterocyclic ring is a five member ring with Y'' = O, one R<sup>5</sup> is -H, two R<sup>5</sup> are -O-(R<sup>2</sup>O)z-H, and at least one R<sup>5</sup> is the following structure -CH(CH<sub>2</sub>-(OR<sup>2</sup>)<sub>z</sub>-H)-CH<sub>2</sub>-(OR<sup>2</sup>)<sub>z</sub>-C(O) R<sup>1</sup> with  $8 \le z + z$ , + z,  $\le 20$  and R<sup>1</sup> is a hydrocarbon with from 8 to 20 carbon atoms and no aryl group;

c. polyhydroxy fatty acid amide surfactants of the formula:

$$R^2 - C(O) - N(R^1) - Z$$

wherein: each  $R^1$  is H,  $C_1$ - $C_4$  hydrocarbyl,  $C_1$ - $C_4$  alkoxyalkyl, or hydroxyalkyl; and  $R^2$  is a  $C_5$ - $C_{31}$  hydrocarbyl moiety; and each Z is a polyhydroxyhydrocarbyl moiety having a linear hydrocarbyl chain with at least 3 hydroxyls directly connected to the chain, or an ethoxylated derivative thereof; and each R' is H or a cyclic mono- or poly- saccharide, or alkoxylated derivative thereof; and

Suitable phase stabilizers also include surfactant complexes formed by one surfactant ion being neutralized with surfactant ion of opposite charge or an electrolyte ion that is suitable for reducing dilution viscosity.

Examples of representative bilayer disrupters include:

## (1)- Alkyl or alkyl-aryl alkoxylated nonionic surfactants

Suitable alkyl alkoxylated nonionic surfactants are generally derived from saturated or unsaturated primary, and secondary fatty alcohols, fatty acids, alkyl phenols, or alkyl aryl (e.g., benzoic) carboxylic acid, where the active hydrogen(s) is alkoxylated with ≤ about 30 alkylene, preferably ethylene, oxide moieties (e.g. ethylene oxide and/or propylene oxide). These nonionic surfactants for use herein preferably have from about 6 to about 22 carbon atoms on the alkyl or alkenyl chain, and are in a straight chain configuration, preferably straight chain configurations having from about 8 to about 18 carbon atoms, with the alkylene oxide being present, preferably at the primary position, in average amounts of ≤ about 30 moles of alkylene oxide per alkyl chain, more preferably from about 3 to about 15 moles of alkylene oxide, and most preferably from about 6 to about 12 moles of alkylene oxide. Preferred materials of this class also have pour points of less than about 70°F (21°C) and/or do not solidify in these softening compositions. Examples of alkyl alkoxylated surfactants with straight chains include Neodol® 91-8, 23-5, 25-9, 1-9, 25-12, 1-9, and 45-13 from Shell, Plurafac® B-26 and C-17 from BASF, and Brij® 76 and 35 from ICI Surfactants. Examples of alkyl-aryl alkoxylated surfactants include: Surfonic N-120 from Huntsman, Igepal® CO-620 and CO-710, from Rhone Poulenc, Triton® N-111 and N-150 from Union Carbide, Dowfax® 9N5 from Dow and Lutensol® AP9 and AP14, from BASF.

#### (2)- Alkyl or alkyl-aryl amine or amine oxide nonionic alkoxylated surfactants

Suitable alkyl alkoxylated nonionic surfactants with amine functionality are generally derived from saturated or unsaturated, primary, and secondary fatty alcohols, fatty acids, fatty methyl esters, alkyl phenol, alkyl benzoates, and alkyl benzoic acids that are converted to amines,

30

5

10

15

20

amine-oxides, and optionally substituted with a second alkyl or alkyl-aryl hydrocarbon with one or two alkylene oxide chains attached at the amine functionality each having ≤ about 50 moles alkylene oxide moieties (e.g. ethylene oxide and/or propylene oxide) per mole of amine. The amine, amide or amine-oxide surfactants for use herein have from about 6 to about 22 carbon atoms, and are in either straight chain or branched chain configuration, preferably there is one hydrocarbon in a straight chain configuration having about 8 to about 18 carbon atoms with one or two alkylene oxide chains attached to the amine moiety, in average amounts of ≤ 50 about moles of alkylene oxide per amine moiety, more preferably from about 3 to about 15 moles of alkylene oxide, and most preferably a single alkylene oxide chain on the amine moiety containing from about 6 to about 12 moles of alkylene oxide per amine moiety. Preferred materials of this class also have pour points less than about 70°F (21°C)and/or do not solidify in these softening compositions. Examples of ethoxylated amine surfactants include Berol® 397 and 303 from Rhone Poulenc and Ethomeens® C/20, C25, T/25, S/20, S/25 and Ethodumeens® T/20 and T25 from Akzo.

Preferably, the compounds of the alkyl or alkyl-aryl alkoxylated surfactants and alkyl or alkyl-aryl amine, amide, and amine-oxide alkoxylated have the following general formula:

$$R_{m}^{1} - Y - [(R^{2}-O)_{z} - H]_{p}$$

wherein each R is selected from the group consisting of saturated or unsaturated, primary, secondary or branched chain alkyl or alkyl-aryl hydrocarbons; said hydrocarbon chain preferably having a length of from about 6 to about 22, more preferably from about 8 to about 18 carbon atoms, and even more preferably from about 8 to about 15 carbon atoms, preferably, linear and with no aryl moiety; wherein each R<sup>2</sup> is selected from the following groups or combinations of the following groups:  $-(CH_2)_n$ - and/or  $-[CH(CH_3)CH_2]$ -; wherein about  $1 < n \le$  about 3; Y is selected from the following groups: -O-; -N(A)<sub>q</sub>-; -C(O)O-; - (O $\leftarrow$ )N(A)<sub>q</sub>-; -B-R<sup>3</sup>-O-; -B-R<sup>3</sup>-N(A)<sub>q</sub>-; - $R^3$ -C(O)O-; -B- $R^3$ -N( $\rightarrow$ O)(A)-; and mixtures thereof; wherein A is selected from the following groups: H; R<sup>1</sup>; -(R<sup>2</sup>-O)<sub>z</sub>-H; -(CH<sub>2</sub>)<sub>x</sub>CH<sub>3</sub>; phenyl, or substituted aryl, wherein  $0 \le x \le$  about 3 and B is selected from the following groups: -O-; -N(A)-; -C(O)O-; and mixtures thereof in which A is as defined above; and wherein each R3 is selected from the following groups:R2; phenyl; or substituted aryl. The terminal hydrogen in each alkoxy chain can be replaced by a short chain C<sub>1-4</sub> alkyl or acyl group to "cap" the alkoxy chain. z is from about 5 to about 30. p is the number of ethoxylate chains, typically one or two, preferably one and m is the number of hydrophobic chains, typically one or two, preferably one and q is a number that completes the structure, usually one.

5

10

15

20

25

Preferred structures are those in which m = 1, p = 1 or 2, and  $5 \le z \le 30$ , and q can be 1 or 0, but when p = 2, q must be 0; more preferred are structures in which m = 1, p = 1 or 2, and  $7 \le z \le 20$ ; and even more preferred are structures in which m = 1, p = 1 or 2, and  $9 \le z \le 12$ . The preferred y is 0.

#### 5 (3)- Alkoxylated and non-alkoxylated nonionic surfactants with bulky head groups

Suitable alkoxylated and non-alkoxylated bilayer disrupters with bulky head groups are generally derived from saturated or unsaturated, primary and secondary fatty alcohols, fatty acids, alkyl phenol, and alkyl benzoic acids that are derivatized with a carbohydrate group or heterocyclic head group. This structure can then be optionally substituted with more alkyl or alkyl-aryl alkoxylated or non-alkoxylated hydrocarbons. The heterocyclic or carbohydrate is alkoxylated with one or more alkylene oxide chains (e.g. ethylene oxide and/or propylene oxide) each having ≤ about 50, preferably ≤ about 30, moles per mole of heterocyclic or carbohydrate. The hydrocarbon groups on the carbohydrate or heterocyclic surfactant for use herein have from about 6 to about 22 carbon atoms, and are in a straight chain configuration, preferably there is one hydrocarbon having from about 8 to about 18 carbon atoms with one or two alkylene oxide chains carbohydrate or heterocyclic moiety with each alkylene oxide chain present in average amounts of ≤ about 50, preferably ≤ about 30, moles of carbohydrate or heterocyclic moiety, more preferably from about 3 to about 15 moles of alkylene oxide per alkylene oxide chain, and most preferably between about 6 and about 12 moles of alkylene oxide total per surfactant molecule including alkylene oxide on both the hydrocarbon chain and on the heterocyclic or carbohydrate moiety. Examples of bilayer disrupters in this class are Tween® 40, 60, and 80 available from ICI Surfactants.

Preferably the compounds of the alkoxylated and non-alkoxylated nonionic surfactants with bulky head groups have the following general formulas:

 $R^{1}-C(O)-Y'-[C(R^{5})]_{m}-CH_{2}O(R_{2}O)_{z}H$ 

wherein  $R^1$  is selected from the group consisting of saturated or unsaturated, primary, secondary or branched chain alkyl or alkyl-aryl hydrocarbons; said hydrocarbon chain having a length of from about 6 to about 22; Y' is selected from the following groups: -O-; -N(A)-; and mixtures thereof; and A is selected from the following groups: H;  $R^1$ ; -( $R^2$ -O)<sub>z</sub>-H; -( $CH_2$ )<sub>x</sub>CH<sub>3</sub>; phenyl, or substituted aryl, wherein  $0 \le x \le$  about 3 and z is from about 5 to about 30; each  $R^2$  is selected from the following groups or combinations of the following groups: -( $CH_2$ )<sub>n</sub>- and/or -

10

15

20

25

5

10

20

25

[CH(CH<sub>3</sub>)CH<sub>2</sub>]-; and each  $R^5$  is selected from the following groups: -OH; and -O( $R^2O$ )<sub>2</sub>-H; and m is from about 2 to about 4;

Another useful general formula for this class of surfactants is

wherein Y" = N or O; and each R<sup>5</sup> is selected independently from the following:

-H, -OH, -(CH<sub>2</sub>)xCH<sub>3</sub>, -(OR<sup>2</sup>)<sub>z</sub>-H, -OR<sup>1</sup>, - OC(O)R<sup>1</sup>, and -CH<sub>2</sub>(CH<sub>2</sub>-(OR<sup>2</sup>)<sub>z</sub>-H)-CH<sub>2</sub>-(OR<sup>2</sup>)<sub>z</sub>-C(O) R<sup>1</sup>. With x R<sup>1</sup>, and R<sup>2</sup>as defined above in section D above and z, z', and z'' are all from about  $5 \le to \le about 20$ , more preferably the total number of z + z' + z'' is from about  $5 \le to \le about 20$ . In a particularly preferred form of this structure the heterocyclic ring is a five member ring with Y'' = O, one R<sup>5</sup> is -H, two R<sup>5</sup> are -O-(R<sup>2</sup>O)<sub>z</sub>-H, and at least one R<sup>5</sup> has the following structure -CH(CH<sub>2</sub>-(OR<sup>2</sup>)<sub>z</sub>-H)-CH<sub>2</sub>-(OR<sup>2</sup>)<sub>z</sub>-OC(O) R<sup>1</sup> with the total z + z' + z'' = to from about  $8 \le to \le about 20$  and R<sup>1</sup> is a hydrocarbon with from about 8 to about 20 carbon atoms and no aryl group.

Another group of surfactants that can be used are polyhydroxy fatty acid amide surfactants of the formula:

$$R^6 - C(O) - N(R^7) - W$$

wherein: each R<sup>7</sup> is H, C<sub>1</sub>-C<sub>4</sub> hydrocarbyl, C<sub>1</sub>-C<sub>4</sub> alkoxyalkyl, or hydroxyalkyl, e.g., 2-hydroxyethyl, 2-hydroxypropyl, etc., preferably C<sub>1</sub>-C<sub>4</sub> alkyl, more preferably C<sub>1</sub> or C<sub>2</sub> alkyl, most preferably C<sub>1</sub> alkyl (i.e., methyl) or methoxyalkyl; and R<sup>6</sup> is a C<sub>5</sub>-C<sub>31</sub> hydrocarbyl moiety, preferably straight chain C<sub>7</sub>-C<sub>19</sub> alkyl or alkenyl, more preferably straight chain C<sub>9</sub>-C<sub>17</sub> alkyl or alkenyl, or mixture thereof; and W is a polyhydroxyhydrocarbyl moiety having a linear hydrocarbyl chain with at least 3 hydroxyls directly connected to the chain, or an alkoxylated derivative (preferably ethoxylated or propoxylated) thereof. W preferably will be derived from a reducing sugar in a reductive amination reaction; more preferably W is a glycityl moiety. W preferably will be selected from the group consisting of -CH<sub>2</sub>-(CHOH)<sub>n</sub>-CH<sub>2</sub>OH, -CH(CH<sub>2</sub>OH)-(CHOH)<sub>n</sub>-CH<sub>2</sub>OH, -CH<sub>2</sub>-(CHOH)<sub>2</sub>(CHOR')(CHOH)-CH<sub>2</sub>OH, where n is an integer from 3 to 5, inclusive, and R' is H or a cyclic mono- or poly- saccharide, and alkoxylated derivatives thereof. Most preferred are glycityls

wherein n is 4, particularly -CH<sub>2</sub>-(CHOH)<sub>4</sub>-CH<sub>2</sub>O. Mixtures of the above W moieties are desirable.

R<sup>6</sup> can be, for example, N-methyl, N-ethyl, N-propyl, N-isopropyl, N-butyl, N-isobutyl, N-2-hydroxyethyl, N-1-methoxypropyl, or N-2-hydroxypropyl.

R<sup>6</sup>-CO-N< can be, for example, cocamide, stearamide, oleamide, lauramide, myristamide, capricamide, palmitamide, tallowamide, etc.

W can be 1-deoxyglucityl, 2-deoxyfructityl, 1-deoxymaltityl, 1-deoxyglactityl, 1-deoxymannityl, 1-deoxymannityl, 1-deoxymaltotriotityl, etc.

### (4)- Alkoxylated cationic quaternary ammonium surfactants

Alkoxylated cationic quaternary ammonium surfactants suitable for this invention are generally derived from fatty alcohols, fatty acids, fatty methyl esters, alkyl substituted phenols, alkyl substituted benzoic acids, and/or alkyl substituted benzoate esters, and/or fatty acids that are converted to amines which can optionally be further reacted with another long chain alkyl or alkyl-aryl group; this amine compound is then alkoxylated with one or two alkylene oxide chains each having ≤ about 50 moles alkylene oxide moieties (e.g. ethylene oxide and/or propylene oxide) per mole of amine. Typical of this class are products obtained from the quaternization of aliphatic saturated or unsaturated, primary, secondary, or branched amines having one or two hydrocarbon chains from about 6 to about 22 carbon atoms alkoxylated with one or two alkylene oxide chains on the amine atom each having less than ≤ about 50 alkylene oxide moieties. The amine hydrocarbons for use herein have from about 6 to about 22 carbon atoms, and are in either straight chain or branched chain configuration, preferably there is one alkyl hydrocarbon group in a straight chain configuration having about 8 to about 18 carbon atoms. Suitable quaternary ammonium surfactants are made with one or two alkylene oxide chains attached to the amine moiety, in average amounts of ≤ about 50 moles of alkylene oxide per alkyl chain, more preferably from about 3 to about 20 moles of alkylene oxide, and most preferably from about 5 to about 12 moles of alkylene oxide per hydrophobic, e.g., alkyl group. Preferred materials of this class also have a pour points below about 70°F (21°C) and/or do not solidify in these softening compositions. Examples of suitable bilayer disrupters of this type include Ethoquad® 18/25, C/25, and O/25 from Akzo and Variquat<sup>®</sup>-66 (soft tallow alkyl bis(polyoxyethyl) ammonium ethyl sulfate with a total of about 16 ethoxy units) from Witco.

Preferably, the compounds of the ammonium alkoxylated cationic surfactants have the following general formula:

5

10

15

20

25

$${R^{1}_{m} - Y - [(R^{2}-O)_{z} - H]_{p}}^{+} X^{-}$$

wherein R<sup>1</sup> and R<sup>2</sup> are as defined previously in section D above;

Y is selected from the following groups: =  $N^+-(A)_q$ ;  $-(CH_2)_n-N^+-(A)_q$ ;  $-B-(CH_2)_n-N^+-(A)_2$ ; -(phenyl)- $N^+-(A)_q$ ; -(B-phenyl)- $N^+-(A)_q$ ; with n being from about 1 to about 4.

Each A is independently selected from the following groups: H;  $R^1$ ;  $-(R^2O)_z$ -H;  $-(CH_2)_x$ CH<sub>3</sub>; phenyl, and substituted aryl; where  $0 \le x \le$  about 3; and B is selected from the following groups: -O-; -NA-; -NA<sub>2</sub>; -C(O)O-; and -C(O)N(A)-; wherein  $R^2$  is defined as hereinbefore; q = 1 or 2; and

X is an anion which is compatible with the softening active ingredient and other components of the softening composition.

Preferred structures are those in which m = 1, p = 1 or 2, and about  $5 \le z \le$  about 50, more preferred are structures in which m = 1, p = 1 or 2, and about  $7 \le z \le$  about 20, and most preferred are structures in which m = 1, p = 1 or 2, and about  $9 \le z \le$  about 12.

### (5)-Alkyl amide alkoxylated nonionic surfactants

Suitable surfactants have the formula:

15

20

25

5

$$R - C(O) - N(R^4)_n - [(R^1O)_x(R^2O)_yR^3]_m$$

wherein R is  $C_{7-21}$  linear alkyl,  $C_{7-21}$  branched alkyl,  $C_{7-21}$  linear alkenyl,  $C_{7-21}$  branched alkenyl, and mixtures thereof. Preferably R is  $C_{8-18}$  linear alkyl or alkenyl.

R<sup>1</sup> is -CH<sub>2</sub>-CH<sub>2</sub>-, R<sub>2</sub> is C<sub>3</sub>-C<sub>4</sub> linear alkyl, C<sub>3</sub>-C<sub>4</sub> branched alkyl, and mixtures thereof; preferably R<sup>2</sup> is -CH(CH<sub>3</sub>)-CH<sub>2</sub>-. Surfactants which comprise a mixture of R1 and R2 units preferably comprise from about 4 to about 12 -CH<sub>2</sub>-CH<sub>2</sub>- units in combination with from about 1 to about 4 -CH(CH<sub>3</sub>)-CH<sub>2</sub>- units. The units may be alternating or grouped together in any combination suitable to the formulator. Preferably the ratio of R<sup>1</sup> units to R<sup>2</sup> units is from about 4:1 to about 8:1. Preferably an R<sup>2</sup> unit (i.e. -C(CH<sub>3</sub>)H-CH<sub>2</sub>-) is attached to the nitrogen atom followed by the balance of the chain comprising from about 4 to 8 -CH<sub>2</sub>-CH<sub>2</sub>- units.

 $R^3$  is hydrogen,  $C_1$ - $C_4$  linear alkyl,  $C_3$ - $C_4$  branched alkyl, and mixtures thereof; preferably hydrogen or methyl, more preferably hydrogen.

R<sup>4</sup> is hydrogen, C<sub>1</sub>-C<sub>4</sub> linear alkyl, C<sub>3</sub>-C<sub>4</sub> branched alkyl, and mixtures thereof; preferably hydrogen. When the index m is equal to 2 the index n must be equal to 0 and the R4 unit is absent.

The index m is 1 or 2, the index n is 0 or 1, provided that m + n equals 2; preferably m is equal to 1 and n is equal to 1, resulting in one -  $[(R^1O)_x(R^2O)_yR^3]$  unit and R4 being present on the nitrogen. The index x is from 0 to about 50, preferably from about 3 to about 25, more preferably from about 3 to about 10. The index y is from 0 to about 10, preferably 0, however when the index y is not equal to 0, y is from 1 to about 4. Preferably all the alkyleneoxy units are ethyleneoxy units.

Examples of suitable ethoxylated alkyl amide surfactants are Rewopal® C<sub>6</sub> from Witco, Amidox® C5 from Stepan, and Ethomid® O / 17 and Ethomid® HT / 60 from Akzo.

#### **High Polymers**

5

10

15

20

25

30

High molecular weight polymers (hereinafter "high polymers") which are substantially compatible with the vehicle can also be useful in order to achieve the desired extensional viscosity characteristics for the softening compositions herein. In one embodiment, the high polymer preferably has a substantially linear chain structure, though a linear chain having short (C<sub>1</sub>-C<sub>3</sub>) branches or a branched chain having one to three long branches are also suitable for use herein. As used herein, the term "substantially compatible" means that the high polymer appears to dissolve in the vehicle as the continuous phase of the softening composition is being prepared (i.e., the continuous phase appears transparent or translucent to the naked eye).

Such polymers also should not destabilize the softening composition due to their presence. For example, a suitable high polymer would not have a sufficiently large number of anionic substituents so as to cause flocculation of the softening composition. It may be necessary to adjust certain properties of the composition in order to insure stability. For example insure that an anionic has a sufficiently low level of anionic character (e.g. via pH adjustment of a presolution of the polymer so as to approach the isoelectric point) so as not to cause flocculation.

Without being bound by theory, it is believed that polymers suitable for use herein preferably self-interact within the vehicle at the molecular level and with particles of the softening active ingredient (e.g. via entanglement, surface absorption, and ionic attraction) in order to increase the extensibility of the softening composition to with a resulting reduction in spray fracture. As used herein the term "spray fracture" is intended to mean separation of the flow of softening composition within a spray apparatus into individual droplets having a size that is sufficiently small that they become aerosolized. The polymers useful herein are preferably high molecular weight, substantially linear chain molecules. The high molecular weight of the polymer enables it to enhance the extensibility of the softening composition such that the composition is suitable for extensional processes in a spray apparatus. It is believed that such spray process

results in droplets or filaments having a size large enough that substantially all of the material deposits onto the web rather than being carried outside the vicinity of the web by air flows adjacent thereto (i.e. the particles are deposited rather than being aerosolized).

As will be recognized, the high polymer should be sufficiently soluble/swellable in the vehicle so as to enable sufficient self-interact and interact with the softener active ingredient particles. As such high polymers forming true solutions in the vehicle or micellar solutions therein are particularly preferred.

In order to effectively interact with other high polymer molecules and with the softening active ingredient particles, the high polymer suitable for use herein should have a weight-average molecular weight of at least 500,000. Typically the weight average molecular weight of the polymer ranges from about 500,000 to about 25,000,000, more typically from about 800,000 to about 22,000,000, even more typically from about 1,000,000 to about 20,000,000, and most typically from about 2,000,000 to about 15,000,000. The high molecular weight polymers are preferred in some embodiments of the invention due to the ability to simultaneously interact with several particles of softening active ingredient, thereby increasing extensional viscosity and reducing spray fracture.

In order to minimize spray fracture, the compositions herein should desirably exhibit certain rheological behavior during spraying, including a certain range of extensional viscosities. Extensional or elongational viscosity ( $\eta_e$ ) relates to the extensibility of the composition, and is particularly important for extensional processes such as spraying. The extensional viscosity includes three types of deformation: uniaxial or simple extensional viscosity, biaxial extensional viscosity, and pure shear extensional viscosity. The uniaxial extensional viscosity is important for uniaxial extensional processes. The other two extensional viscosities are important for the biaxial extension or forming processes such as droplet formation. Without being bound by theory, it is believed that the "stringiness" caused by increased extensional viscosity results in filament formation and an increased droplet size during spraying, for a given air pressure, because the increased viscosity provides resistance to the force exerted on the nascent droplet by the air jets within the spray nozzle allowing additional material to pass through the fluid orifice before a droplet separates from the fluid stream.

It has been found that softening compositions useful for spray application according to the present invention typically have their extensional viscosity increased by a factor of at least 2 when a selected high polymer is added to the composition, preferably by a factor of at least about 5. Preferably, the compositions of present invention show an increase in the extensional viscosity

5

10

. 15

20

25

of a factor of about 5 to about 500, more preferably of about 20 to about 300, and most preferably from about 30 to about 100, when a selected high polymer is added. The higher the level of the high polymer, the greater the increase in extensional viscosity. The higher the softening active ingredient solids content at a given high polymer concentration, the higher the extensional viscosity. The extensional viscosity of the compositions of the present invention is at least about 5 pascal • seconds at a Hencky strain of 6. In one embodiment of the present invention, the softening compositions of the present invention further have an extensional viscosity in the range of from about 5 pascal • seconds to about 5,000 pascal • seconds, typically from about 5 pascal • seconds to about 750 pascal • seconds, even more typically from about 5 pascal • seconds to about 600 pascal • seconds and most typically from about 5 pascal • seconds to about 100 pascal • seconds at the die temperature. The extensional viscosity is calculated according to the method set forth hereinafter in the Analytical Methods section.

5

10

15

20

25

30

The Trouton ratio (Tr) is often used to express the extensional flow behavior. The Trouton ratio is defined as the ratio between the extensional viscosity ( $\eta_e$ ) and the shear viscosity ( $\eta_s$ ),

$$Tr = \eta_e(\varepsilon^{\bullet}, t) / \eta_s$$

wherein the extensional viscosity  $\eta_e$  is dependent on the deformation rate ( $\epsilon^{\bullet}$ ) and time (t). For a Newtonian fluid, the uniaxial extension Trouton ratio has a constant value of 3. For a non-Newtonian fluid, such as the compositions herein, the extensional viscosity is dependent on the deformation rate ( $\epsilon^{\bullet}$ ) and time (t). It has also been found that compositions of the present invention typically have a Trouton ratio of at least about 3. Typically, the Trouton ratio ranges from about 10 to about 5,000, more typically from about 20 to about 1,000, even more typically from about 30 to about 500, when measured at the processing temperature and 700 s<sup>-1</sup>. As used herein "Processing Temperature" means the temperature of the composition when the composition is applied to the tissue web.

The increased extensional viscosity also becomes apparent when shear viscosity is measured at different shear rates. Specifically for softening compositions according to the present invention that comprise a high polymer it has been found that low shear viscosity (~10 sec<sup>-1</sup>) increases by a very large amount, compared with a composition without the high polymer, but the high shear (>100 sec<sup>-1</sup>) is not affected very much.

Nonlimiting examples of suitable high polymers include polyacrylamide and certain derivatives acrylic polymers and copolymers as may be compatible with the softening composition of the present invention; vinyl polymers including polyvinyl alcohol; polyvinylacetate; polyvinylpyrrolidone; polyethylene vinyl acetate; polyethyleneimine; and the like; polyalkylene oxides such as polyethylene oxide; polypropylene oxide; polyethylene/propylene oxide; and mixtures thereof. Copolymers made from mixtures of monomers selected from any of the aforementioned polymers are also suitable herein. Other exemplary high polymers include water soluble polysaccharides such as alginates, carrageenans, pectin and derivatives, chitin and derivatives, and the like; gums such as guar gum, xanthum gum, agar, gum arabic, karaya gum, tragacanth gum, locust bean gum, and like gums; water soluble derivatives of cellulose, such as alkylcellulose, hydroxyalkylcellulose, carboxyalkylcellulose, and the like; and mixtures thereof.

Some polymers (e.g., polyacrylic acid, polymethacrylic acid) are generally not available in the high molecular weight range (i.e., 500,000 or higher). A small amount of crosslinking agents may be added to create branched polymers of suitably high molecular weight useful herein.

The high polymer, when used in a spraying process, is added to the composition of the present invention in an amount effective to visibly reduce spray fracture and the resulting aerosolization during the spraying process such that substantially all of the softening composition is deposited onto the tissue web. These polymers, when used, are typically present in the range from about 0.01 to about 5 wt%, more typically from about 0.01 to about 2 wt%, even more typically from about 0.01 to about 1 wt%, and most typically from about 0.05 to about 0.5 wt% of the composition. A particularly preferred range is between about 0.1 wt% and about 0.25 wt%. It is surprising to find that at a relatively low concentration, these polymers can significantly improve the air pressure operating window in a spray apparatus.

The following is a nonlimiting list of materials suitable for use as a high polymer for purposes of the present invention. Superfloc® N-300 is a polyacrylamide having a weight-average molecular weight of about 10,000,000 and Superfloc N-300LMW with a weight average molecular weight of about 5,000,000 both of which are available from Cytec Co., Stamford, CT. Nonionic polyacrylamides PAM-a and PAM-b having a weight-average molecular weight of 15,000,000, and 5,000,000 to 6,000,000, respectively, are available from Scientific Polymer Products, Inc., Ontario, NY. Polyethyleneimine having a weight-average molecular weight of 750,000 is available from Aldrich Chemical Co., Milwaukee, WI. High molecular weight cellulose as is available from Hercules of Wilmington, DE as Aqualon® cellulose gum.

5

10

15

20

25

## Minor Components of the Softening Composition

The vehicle can also comprise minor ingredients as may be known to the art. examples include: mineral acids or buffer systems for pH adjustment (may be required to maintain hydrolytic stability for certain softening active ingredients) and antifoam ingredients (e. g., a silicone emulsion as is available from Dow Corning, Corp. of Midland, MI as Dow Corning 2310) as a processing aid to reduce foaming when the softening composition of the present invention is applied to a web of tissue.

It may also be desirable to provide means to control the activity of undesirable microorganisms in the softening composition of the present invention. It is known that organisms, such as bacteria, molds, yeasts, and the like, can cause degradation of the composition on storage. Undesirable organisms can also potentially transfer to users of tissue paper products that are softened with a composition according to the present invention that is contaminated by such organisms. These undesirable organisms can be controlled by adding an effective amount of a biocidal material to the softening composition. Proxel GXL, as is available from Avecia, Inc. of Wilmington, DE, has been found to be an effective biocide in the composition of the present invention when used at a level of about 0.1%. Alternatively, the pH of the composition can be made more acid to create a more hostile environment for undesirable microorganisms. Means such as those described above can be used to adjust the pH to be in a range of between about 2.5 to 4.0, preferably between about 2.5 and 3.5, more preferably between about 2.5 and about 3.0 so as to create such a hostile environment.

Stabilizers may also be used to improve the uniformity and shelf life of the dispersion. For example, an ethoxylated polyester, HOE S 4060, available from Clariant Corporation of Charlotte, NC may be included for this purpose.

Process aids may also be used, including for example, a brightener, such as Tinopal CBS-X, obtainable from CIBA-GEIGY of Greensboro, NC may be added to the dispersion to allow easy qualitative viewing of the application uniformity, via inspection of the finished tissue web, containing a surface-applied softening composition, under UV light.

#### Forming the Softening Composition

As noted above, the softening composition of the present invention is a dispersion of a softening active ingredient in a vehicle. Depending on the softening active ingredient chosen, the desired application level and other factors as may require a particular level of softening active ingredient in the composition, the level of softening active ingredient may vary between about

5

10

15

20

25

10% of the composition and about 50% of the composition. Preferably, the softening active ingredient comprises between about 25% and about 45% of the composition. Most preferably, the softening active ingredient comprises between about 30% and about 40% of the composition. The nonionic surfactant is present at a level between about 1% and about 15% of the level of the softening active ingredient, preferably between about 2% and about 10%. The composition further comprises the high polymer at a level of between about 0.01% and about 5%. Depending on the method used to produce the softening active ingredient the softening composition may also comprise between about 2% and about 30%, preferably between about 5% and about 25% of a plasticizer. As noted above, the preferred primary component of the vehicle is water. In addition, the vehicle preferably comprises an alkali or alkaline earth salt of a simple organic acid electrolyte and may comprise minor ingredients to adjust pH, to control foam, or to aid in stability of the dispersion. The following describes preparation of a particularly preferred softening composition of the present invention.

A particularly preferred softening composition of the present invention is prepared as follows. The materials comprising this composition are more specifically defined in Table 1 which follows this description. Amounts used in each step are sufficient to result in the finished composition detailed in that table. The appropriate quantity of water is heated (extra water may be added to compensate for evaporation loss) to about 165°F (75°C). The high polymer, sulfuric acid (38% solution) and antifoam ingredient are added. Concurrently, the blend of softening active ingredient and plasticizer is melted by heating it to a temperature of about 150°F (65°C). The melted mixture of softening active ingredient, plasticizer, and nonionic surfactant is then slowly added to the heated acidic aqueous phase with mixing to evenly distribute the disperse phase throughout the vehicle. (The water solubility of the polyethylene glycol probably carries it into the continuous phase, but this is not essential to the invention and plasticizers which are more hydrophobic and thus remain associated with the alkyl chains of the quaternary ammonium compound are also allowed within the scope of the present invention.) Once the softening active ingredient is thoroughly dispersed, part of the sodium formate is added (as a 5% solution) intermittently with mixing to provide an initial viscosity reduction. The stabilizer is then slowly added to the mixture with continued agitation. The remainder of the sodium formate (as a 25% solution) is then added. Lastly, nonionic surfactant is added with continued mixing.

Table 1

5

10

15

20

25

Component	Concentration
Continuous Phase	
Water	QS to 100%
Electrolyte <sup>1</sup>	2.0%
Antifoam <sup>2</sup>	0.25%
Bilayer Disrupter <sup>3</sup>	0.8%
Sulfuric Acid⁴	1.1%
Plasticizer <sup>5</sup>	20.0%
Stabilizer <sup>6</sup>	1.4%
High polymer <sup>7</sup>	0.1%
Disperse Phase	
	45.004

Softening Active Ingredient<sup>5</sup> 45.0%

1. 0.25 % from 5 % aqueous sodium formate solution and 1.75 % from 25 % aqueous sodium formate solution

2. Silicone Emulsion (10% active)—Dow Corning 2310®, marketed by Dow Corning Corp., Midland, MI

3. Suitable nonionic surfactants are available from Shell Chemical of Houston, TX under the trade name NEODOL 91-8.

4. Available as a 38% solution from J. T. Baker Chemical Company of Phillipsburg, NJ

5. Plasticizer and softening active ingredient obtained pre-blended from Goldschmidt Chemical Company of Dublin OH as DXP5558-6 and comprises about 31 parts polyethylene glycol 400 and about 69 parts tallow diester quaternary)

6. Stabilizer is HOE S 4060, from Clariant Corp., Charlotte, NC

7. High polymer is a nonionic polyacrylamide Superfloc N-300 as is available from Cytec Co., Stamford, CT

The resulting chemical softening composition is a milky, low viscosity dispersion suitable for application to cellulosic structures as described below for providing desirable tactile softness to such structures. It displays a shear-thinning non-Newtonian viscosity. Suitably, the composition has a viscosity less than about 1000 centipoise (cp), as measured at 25° C and at a shear rate of 100 sec<sup>-1</sup> using the method described in the TEST METHODS section below. Preferably, the composition has a viscosity less than about 500 cp. More preferably, the viscosity is less than about 300 cp.

#### Application Method

In one preferred embodiment, the softening composition of the current invention may be applied after the tissue web has been dried and creped, and, more preferably, while the web is still at an elevated temperature. Preferably, the softening composition is applied to the dried and

creped tissue web before the web is wound onto the parent roll. Thus, in a preferred embodiment of the present invention the softening composition is applied to a hot, overdried tissue web after the web has been creped and after the web has passed through the calender rolls which control the caliper. Still more preferably, the composition is applied only to a side of the web that does not contact any rolls between the calendar rolls and the winder.

The softening composition described above is preferably applied to the web in a macroscopically uniform fashion so that substantially the entire sheet benefits from the effect of the softening composition. Following application to the hot web, at least a portion of the volatile components of the vehicle preferably evaporates leaving preferably a thin film containing any remaining unevaporated portion of the volatile components of the vehicle, the softening active ingredient, and other nonvolatile components of the softening composition. By "thin film" is meant any thin coating, haze or mist on the on the web. This thin film can be microscopically continuous or be comprised of discrete elements. If the thin film is comprised of discrete elements, the elements can be of uniform size or varying in size; further they may be arranged in a regular pattern or in an irregular pattern, but macroscopically the thin film is uniform. Preferably the thin film is composed of discrete elements.

The softening composition can be added to either side of the tissue web singularly, or to both sides. Preferably, the composition is applied only to a side of the web that does not contact any rolls between the calendar rolls and the parent roll.

A preferred method of macroscopically uniformly applying the softening composition to the web is spraying. Spraying has been found to be economical, and can be accurately controlled with respect to quantity and distribution of the softening composition, so it is more preferred. The dispersed softening composition is applied onto the dried, creped tissue web after the Yankee dryer and before the parent roll. A particularly convenient means of accomplishing this application is to apply the softening composition to the web after the calendar rolls and before the parent roll. A particularly preferred application position is between the calendar rolls and any spreading roll that may be positioned between the calendar rolls and the parent roll. Such position is particularly preferred because the web is controlled by rolls at each end of the span where the composition is applied and there is still some web path length before the web is wound onto the parent roll for volatilization of the vehicle.

Figure 1 illustrates a preferred method of applying the softening composition to the tissue web. Referring to Figure 1, a wet tissue web 1 is on carrier fabric 14 past turning roll 2 and transferred to Yankee dryer 5 by the action of pressure roll 3 while carrier fabric 14 travels past

5

10

20

25

5

10

15

20

25

30

turning roll 16. The web is adhesively secured to the cylindrical surface of Yankee dryer 5 by adhesive applied by spray applicator 4. Drying is completed by steam-heated Yankee dryer 5 and by hot air which is heated and circulated through drying hood 6 by means not shown. The web is then dry creped from the Yankee dryer 5 by doctor blade 7, after which it is designated creped paper sheet 15. Paper sheet 15 then passes through calendar rolls 10 and 11. The softening composition is then applied to sheet 15 by spray applicator 8 in the span between calendar rolls 10, 11 and spreading roll 9. The treated sheet 15 then travels over a circumferential portion of reel 12 and is wound onto parent roll 13 after a portion of the vehicle has evaporated as the web passes through the span between spreading roll 9 and reel 12.

Equipment suitable for spraying softening composition of the present invention onto the tissue web of the present invention include external mix, air atomizing nozzles, such as are available from ITW-Dynatec of Hendersonville, TN as UFD spray tips.

Suitably, the softening composition is disposed at a level of between about 0.1% and about 8% of the weight of the paper sheet 15, preferably between about 0.1% and about 5%, more preferably between about 0.1% and about 3%.

While not wishing to be bound by theory or to otherwise limit the present invention, the following description of typical process conditions encountered during the papermaking operation and their impact on the process described in this invention is provided. The Yankee dryer raises the temperature of the tissue sheet and removes the moisture. The steam pressure in the Yankee is on the order of 110 PSI (750 kPa). This pressure is sufficient to increase the temperature of the cylinder to about 170°C. The temperature of the paper on the cylinder is raised as the water in the sheet is removed. The temperature of the sheet as it leaves the doctor blade can be in excess of 120°C. The sheet travels through space to the calender and the reel and loses some of this heat. The temperature of the paper wound in the reel is measured to be on the order of 60°C. Eventually the sheet of paper cools to room temperature. This can take anywhere from hours to days depending on the size of the paper roll. As the paper cools it also absorbs moisture from the atmosphere.

Since the softening composition of the present invention is applied to the paper while it is overdried, the water added to the paper with the softening composition by this method (i.e. residual water that does not evaporate in the span between spreading roll 9 and reel 12) is not sufficient to cause the paper to lose a significant amount of its strength and thickness. Thus, no further drying is required.

#### Example 1

5

This Example illustrates preparation of tissue paper exhibiting one embodiment of the present invention. This example demonstrates the production of homogeneous tissue paper webs that are provided with a preferred embodiment of the softening composition of the present invention made as described above. The composition is applied to one side of the web and the webs are combined into a two-ply bath tissue product.

A pilot scale Fourdrinier papermaking machine is used in the practice of the present invention.

An aqueous slurry of NSK of about 3% consistency is made up using a conventional repulper and is passed through a stock pipe toward the headbox of the Fourdrinier.

In order to impart temporary wet strength to the finished product, a 1% dispersion of Parez 750<sup>®</sup> is prepared and is added to the NSK stock pipe at a rate sufficient to deliver 0.3% Parez 750<sup>®</sup> based on the dry weight of the NSK fibers. The absorption of the temporary wet strength resin is enhanced by passing the treated slurry through an in-line mixer.

An aqueous slurry of eucalyptus fibers of about 3% by weight is made up using a conventional repulper. The stock pipe carrying eucalyptus fibers is treated with a cationic starch, RediBOND 5320®, which is delivered as a 2% dispersion in water and at a rate of 0.15% based on the dry weight of starch and the finished dry weight of the resultant creped tissue product. Absorption of the cationic starch is improved by passing the resultant mixture through an in line mixer.

The stream of NSK fibers and eucalyptus fibers are then combined in a single stock pipe prior to the inlet of the fan pump. The combined NSK fibers and eucalyptus fibers are then diluted with white water at the inlet of a fan pump to a consistency of about 0.2% based on the total weight of the NSK fibers and eucalyptus fibers.

The homogeneous slurry of NSK fibers and eucalyptus fibers are directed into a multichanneled headbox suitably equipped to maintain the homogeneous stream until discharged onto a traveling Fourdrinier wire. The homogeneous slurry is discharged onto the traveling Fourdrinier wire and is dewatered through the Fourdrinier wire and is assisted by a deflector and vacuum boxes.

The embryonic wet web is transferred from the Fourdrinier wire, at a fiber consistency of about 15% at the point of transfer, to a patterned drying fabric. The drying fabric is designed to yield a pattern densified tissue with discontinuous low-density deflected areas arranged within a

5

10

15

20

25

30

continuous network of high density (knuckle) areas. This drying fabric is formed by casting an impervious resin surface onto a fiber mesh supporting fabric. The supporting fabric is a 45 x 52 filament, dual layer mesh. The thickness of the resin cast is about 10 mil above the supporting fabric. The knuckle area is about 40% and the open cells remain at a frequency of about 562 per square inch.

Further dewatering is accomplished by vacuum assisted drainage until the web has a fiber consistency of about 28%.

While remaining in contact with the patterned forming fabric, the patterned web is pre-dried by air blow-through predryers to a fiber consistency of about 62% by weight.

The semi-dry web is then transferred to the Yankee dryer and adhered to the surface of the Yankee dryer with a sprayed creping adhesive comprising a 0.125% aqueous solution of polyvinyl alcohol. The creping adhesive is delivered to the Yankee surface at a rate of 0.1% adhesive solids based on the dry weight of the web.

The fiber consistency is increased to about 96% before the web is dry creped from the Yankee with a doctor blade.

The doctor blade has a bevel angle of about 25 degrees and is positioned with respect to the Yankee dryer to provide an impact angle of about 81 degrees. The Yankee dryer is operated at a temperature of about 350°F (177°C) and a speed of about 800 fpm (feet per minute) (about 244 meters per minute).

The web is then passed between two calender rolls. The two calender rolls are biased together at roll weight and operated at surface speeds of 656 fpm (about 200 meters per minute) which produces a percent crepe of about 18%.

At a location after the calendar rolls, the web is sprayed with a chemical softening composition, further described below, using the aforementioned UFD nozzle. The composition is sprayed on the surface opposite to that contacted by the downstream spreading roll.

Materials used in the preparation of the chemical softening mixture are:

1. Partially hydrogenated tallow diester chloride quaternary ammonium compound premixed with polyethylene glycol 400. The premix is 70% quaternary ammonium compound (Adogen SDMC-type from Witco incorporated and 30% PEG 400, available from J.T. Baker Company of Phillipsburg, NJ) as DXP-505-91.

2. Neodol 91-8, an ethoxylated fatty alcohol from Shell chemical of Houston, TX.

- 3. Calcium chloride pellets from J. T. Baker, Company of Phillipsburg, NJ.
- Polydimethylsiloxane 10 percent dispersion in water (DC2310) from Dow Corning of Midland, MI.
- 5. Sulfuric acid from J. T. Baker Company of Phillipsburg, NJ.
  - 6. Brightener is Tinopal CBS-X, obtainable from CIBA-GEIGY of Greensboro, NC.
  - 7. Stabilizer is HOE S 4060, from Clariant Corp., Charlotte, NC.
  - 8. Polyacrylamide (Superfloc N-300 as is available from Cytec Co., Stamford, CT).
- These materials are prepared as follows to form the softening composition of the present invention.

The chemical softening composition (Composition 1) is prepared by heating the required quantity of water to about 75°C and adding the high polymer (Superfloc N-300), nonionic surfactant (Neodol 91-8), the brightener, and the polydimethylsiloxane to the heated water. The solution is then adjusted to a pH of about 4 using hydrochloric acid. The premix of quaternary compound and PEG 400 is then heated to about 65°C and metered into the water premix with stirring until the mixture is fully homogeneous. About half of the calcium chloride is added as a 2.5% solution in water with continued stirring. The stabilizer is then added with continued mixing. Final viscosity reduction is achieved by adding the remainder of the calcium chloride (as a 25% solution) with continued mixing. The components are blended in a proportion sufficient to provide a composition having the following approximate concentrations:

	35%	Partially hydrogenated tallow diester chloride quaternary ammonium compound
• .	47%	Water
25	15%	PEG 400
	0.9%	Neodol 23-5
	0.25%	Superfloc N-300
	0.5%	CaCl <sub>2</sub>
	1.2%	Stabilizer
30	0.15%	Polydimethylsiloxane

15

0.13% HCl

0.1% Brightener

After cooling, the composition has a viscosity of about 450 cp as measured at 25° C and at a shear rate of 100 sec-1 using the method described in the TEST METHODS section.

The chemical softening composition is sprayed onto the web downstream of the calendar rolls. The resulting tissue paper has a basis weight of about 12.8 lb per 3000 ft<sup>2</sup>.

The web is converted into a homogeneous, creped patterned densified tissue paper product. The resulting treated tissue paper has an improved tactile sense of softness relative to the untreated control.

#### 10 Example 2

5

This Example illustrates the effect of adding a high polymer on extensional viscosity. A composition substantially the same as that described in Example 1 is prepared (Composition 2). Significant compositional differences are:

- 1) The high polymer is removed from the composition.
- 2) The level of softening active ingredient is increased from about 35% to about 40%.
- 3) The level of plasticized is increased from about 15% to about 17%.

#### Example 3

Compositions 1 and 2 are evaluated for extensional viscosity at a Hencky strain of 6 using the method described in the TEST METHODS section with the following results.

20

15

Table 2

Composition	Approximate Extensional Viscosity
	(pascal • seconds)
1	<0.1
2	375

#### TEST METHODS

#### Softening Active Ingredient Level on Tissue

Analysis of the amounts of softening active ingredients described herein that are retained on tissue paper webs can be performed by any method accepted in the applicable art. These methods

are exemplary, and are not meant to exclude other methods which may be useful for determining levels of particular components retained by the tissue paper.

The following method is appropriate for determining the quantity of the preferred quaternary ammonium compounds (QAC) that may deposited by the method of the present invention. A standard anionic surfactant (sodium dodecylsulfate—NaDDS) solution is used to titrate the QAC using a dimidium bromide indicator.

# Preparation of Standard Solutions

The following methods are applicable for the preparation of the standard solutions used in this titration method.

# 10 Preparation of Dimidium Bromide Indicator

To a 1 liter volumetric flask:

- A) Add 500 milliliters of distilled water.
- B) Add 40 ml. of dimidium bromide-disulphine blue indicator stock solution, available from Gallard-Schlesinger Industries, Inc. of Carle Place, NY.
- 15 C) Add 46 ml. of 5N H<sub>2</sub>SO<sub>4</sub>
  - Fill flask to the mark with distilled water and mix.

Preparation of the NaDDS solution. to a 1 liter volumetric flask:

- A) Weigh 0.1154 grams of NaDDS available from Aldrich Chemical Co. of Milwaukee, WI as sodium dodecyl sulfate (ultra pure).
- 20 B) Fill flask to mark with distilled water and mix to form a 0.0004N solution.

### **Method**

- On an analytical balance, weigh approximately 0.5 grams of tissue. Record the sample weight to the nearest 0.1 mg.
- Place the sample in a glass cylinder having a volume of about 150 milliliters which contains
   a star magnetic stirrer. Using a graduated cylinder, add 20 milliliters of methylene chloride.
  - 3. In a fume hood, place the cylinder on a hot plate turned to low heat. Bring the solvent to a full boil while stirring and using a graduated cylinder, add 35 milliliters of dimidium bromide indicator solution.

4. While stirring at high speed, bring the methylene chloride to a full boil again. Turn off the heat, but continue to stir the sample. The QAC will complex with the indicator forming a blue colored compound in the methylene chloride layer.

- 5. Using a 10 ml. burette, titrate the sample with a solution of the anionic surfactant. This is done by adding an aliquot of titrant and rapidly stirring for 30 seconds. Turn off the stir plate, allow the layers to separate, and check the intensity of the blue color. If the color is dark blue add about 0.3 milliliters of titrant, rapidly stir for 30 seconds and turn off stirrer. Again check the intensity of the blue color. Repeat if necessary with another 0.3 milliliters When the blue color starts to become very faint, add the titrant dropwise between stirrings. The endpoint is the first sign of a slight pink color in the methylene chloride layer.
  - 6. Record the volume of titrant used to the nearest 0.05 ml.
  - 7. Calculate the amount of QAC in the product using the equation:

$$\frac{(millilitersNaDDS - X) \times Y \times 2}{SampleWt(Grams)} = PoundsPerTonQAC$$

Where X is a blank correction obtained by titrating a specimen without the QAC of the present invention. Y is the milligrams of QAC that 1.00 milliliters of NaDDS will titrate. (For example, Y=0.254 for one particularly preferred QAC, i.e. diestherdi(touch-hydrogenated)tallow dimethyl chloride.)

#### Tissue Density

The density of tissue paper, as that term is used herein, is the average density calculated as the basis weight of that paper divided by the caliper, with the appropriate unit conversions incorporated therein. Caliper of the tissue paper, as used herein, is the thickness of the paper when subjected to a compressive load of 95 g/in<sup>2</sup> (15.5 g/cm<sup>2</sup>).

#### **Extensional Viscosity**

The extensional viscosity is measured using a capillary thinning rheometer as described in Bazilevskii, et al, <u>Polymer Science Series A</u>, "Failure of Polymer Solution Filaments", vol. 39, 3 (1997), pp 316-324. In summary, extensional viscosity is measured by introducing a sample between two plates, rapidly separating the plates, and measuring filament diameter as the plates separate. Meaningful differences between the disclosure of Brazilevski, et al. and the instrument used herein include:

1) The actuator is based on two air pistons.

30

5

10

15

- 2) Both the top and bottom plates are tracked.
- 3) Filament diameter is measured by a scanning laser micrometer rather than light intensity measurement.
- 4) End plate diameter is 4.9 +/- 0.1 mm.
- 5) Initial end plate gap is 3-5 mm.

5

15

20

25

30

6) Final end plate gap is 10-12 mm.

Extensional viscosity is measured at Hencky Strain of about 6.

## Panel Softness of Tissue Papers

Ideally, prior to softness testing, the paper samples to be tested should be conditioned according to TAPPI Method #T402OM-88. Preferably, samples are preconditioned for 24 hours at 10 to 35% relative humidity and within a temperature range of 22 to 40°C. After this preconditioning step, samples should be conditioned for 24 hours at a relative humidity of 48 to 52% and within a temperature range of 22 to 24°C.

Ideally, the softness panel testing should take place within the confines of a constant temperature and humidity room. If this is not feasible, all samples, including the controls, should experience identical environmental exposure conditions.

Softness testing is performed as a paired comparison in a form similar to that described in "Manual on Sensory Testing Methods", ASTM Special Technical Publication 434, published by the American Society For Testing and Materials 1968 and is incorporated herein by reference. Softness is evaluated by subjective testing using what is referred to as a Paired Difference Test. The method employs a standard external to the test material itself. For tactilely perceived softness two samples are presented such that the subject cannot see the samples, and the subject is required to choose one of them on the basis of tactile softness. The result of the test is reported in what is referred to as Panel Score Unit (PSU). With respect to softness testing to obtain the softness data reported herein in PSU, a number of softness panel tests are performed. In each test ten practiced softness judges are asked to rate the relative softness of three sets of paired samples. The pairs of samples are judged one pair at a time by each judge: one sample of each pair being designated X and the other Y. Briefly, each X sample is graded against its paired Y sample as follows:

1. a grade of plus one is given if X is judged to may be a little softer than Y, and a grade of minus one is given if Y is judged to may be a little softer than X;

2. a grade of plus two is given if X is judged to surely be a little softer than Y, and a grade of minus two is given if Y is judged to surely be a little softer than X;

- 3. a grade of plus three is given to X if it is judged to be a lot softer than Y, and a grade of minus three is given if Y is judged to be a lot softer than X; and, lastly:
- 4. a grade of plus four is given to X if it is judged to be a whole lot softer than Y, and a grade of minus 4 is given if Y is judged to be a whole lot softer than X.

The grades are averaged and the resultant value is in units of PSU. The resulting data are considered the results of one panel test. If more than one sample pair is evaluated then all sample pairs are rank ordered according to their grades by paired statistical analysis. Then, the rank is shifted up or down in value as required to give a zero PSU value to which ever sample is chosen to be the zero-base standard. The other samples then have plus or minus values as determined by their relative grades with respect to the zero base standard. The number of panel tests performed and averaged is such that about 0.2 PSU represents a significant difference in subjectively perceived softness.

### Strength of Tissue Papers

#### Dry Tensile Strength

This method is intended for use on finished paper products, reel samples, and unconverted stocks. The tensile strength of such products may be determined on one inch wide strips of sample using a Thwing-Albert Intelect II Standard Tensile Tester (Thwing-Albert Instrument Co of Philadelphia, PA).

#### Sample Conditioning and Preparation

Prior to tensile testing, the paper samples to be tested should be conditioned according to TAPPI Method #T402OM-88. All plastic and paper board packaging materials must be carefully removed from the paper samples prior to testing. The paper samples should be conditioned for at least 2 hours at a relative humidity of 48 to 52% and within a temperature range of 22 to 24 °C. Sample preparation and all aspects of the tensile testing should also take place within the confines of the constant temperature and humidity room.

For finished product, discard any damaged product. Next, remove 5 strips of four usable units (also termed sheets) and stack one on top to the other to form a long stack with the perforations between the sheets coincident. Identify sheets 1 and 3 for machine direction tensile

5

10

15

20

25

measurements and sheets 2 and 4 for cross direction tensile measurements. Next, cut through the perforation line using a paper cutter (JDC-1-10 or JDC-1-12 with safety shield from Thwing-Albert Instrument Co. of Philadelphia, PA) to make 4 separate stocks. Make sure stacks 1 and 3 are still identified for machine direction testing and stacks 2 and 4 are identified for cross direction testing.

Cut two 1" wide strips in the machine direction from stacks 1 and 3. Cut two 1" wide strips in the cross direction from stacks 2 and 4. There are now four 1" wide strips for machine direction tensile testing and four 1" wide strips for cross direction tensile testing. For these finished product samples, all eight 1" wide strips are five usable units (also termed sheets) thick.

For unconverted stock and/or reel samples, cut a 15" by 15" sample which is 8 plies thick from a region of interest of the sample using a paper cutter (JDC-1-10 or JDC-1-12 with safety shield from Thwing-Albert Instrument Co of Philadelphia, PA). Make sure one 15" cut runs parallel to the machine direction while the other runs parallel to the cross direction. Make sure the sample is conditioned for at least 2 hours at a relative humidity of 48 to 52% and within a temperature range of 22 to 24 °C. Sample preparation and all aspects of the tensile testing should also take place within the confines of the constant temperature and humidity room.

From this preconditioned 15" by 15" sample which is 8 plies thick, cut four strips 1" by 7" with the long 7" dimension running parallel to the machine direction. Note these samples as machine direction reel or unconverted stock samples. Cut an additional four strips 1" by 7" with the long 7" dimension running parallel to the cross direction. Note these samples as cross direction reel or unconverted stock samples. Make sure all previous cuts are made using a paper cutter (JDC-1-10 or JDC-1-12 with safety shield from Thwing-Albert Instrument Co. of Philadelphia, PA). There are now a total of eight samples: four 1" by 7" strips which are 8 plies thick with the 7" dimension running parallel to the machine direction and four 1" by 7" strips which are 8 plies thick with the 7" dimension running parallel to the cross direction.

#### Operation of Tensile Tester

For the actual measurement of the tensile strength, use a Thwing-Albert Intelect II Standard Tensile Tester (Thwing-Albert Instrument Co. of Philadelphia, PA). Insert the flat face clamps into the unit and calibrate the tester according to the instructions given in the operation manual of the Thwing-Albert Intelect II. Set the instrument crosshead speed to 4.00 in/min and the 1st and 2nd gauge lengths to 2.00 inches. The break sensitivity should be set to 20.0 grams and the sample width should be set to 1.00" and the sample thickness at 0.025".

5

10

15

20

25

A load cell is selected such that the predicted tensile result for the sample to be tested lies between 25% and 75% of the range in use. For example, a 5000 gram load cell may be used for samples with a predicted tensile range of 1250 grams (25% of 5000 grams) and 3750 grams (75% of 5000 grams). The tensile tester can also be set up in the 10% range with the 5000 gram load cell such that samples with predicted tensiles of 125 grams to 375 grams could be tested.

Take one of the tensile strips and place one end of it in one clamp of the tensile tester. Place the other end of the paper strip in the other clamp. Make sure the long dimension of the strip is running parallel to the sides of the tensile tester. Also make sure the strips are not overhanging to the either side of the two clamps. In addition, the pressure of each of the clamps must be in full contact with the paper sample.

After inserting the paper test strip into the two clamps, the instrument tension can be monitored. If it shows a value of 5 grams or more, the sample is too taut. Conversely, if a period of 2-3 seconds passes after starting the test before any value is recorded, the tensile strip is too slack.

Start the tensile tester as described in the tensile tester instrument manual. The test is complete after the crosshead automatically returns to its initial starting position. Read and record the tensile load in units of grams from the instrument scale or the digital panel meter to the nearest unit.

If the reset condition is not performed automatically by the instrument, perform the necessary adjustment to set the instrument clamps to their initial starting positions. Insert the next paper strip into the two clamps as described above and obtain a tensile reading in units of grams. Obtain tensile readings from all the paper test strips. It should be noted that readings should be rejected if the strip slips or breaks in or at the edge of the clamps while performing the test.

#### Calculations

5

10

15

20

25

30

For the four machine direction 1" wide finished product strips, sum the four individual recorded tensile readings. Divide this sum by the number of strips tested. This number should normally be four. Also divide the sum of recorded tensiles by the number of usable units per tensile strip. This is normally five for both 1-ply and 2-ply products.

Repeat this calculation for the cross direction finished product strips.

For the unconverted stock or reel samples cut in the machine direction, sum the four individual recorded tensile readings. Divide this sum by the number of strips tested. This number

should normally be four. Also divide the sum of recorded tensiles by the number of usable units per tensile strip. This is normally eight.

Repeat this calculation for the cross direction unconverted or reel sample paper strips.

All results are in units of grams/inch.

For purposes of this specification, the tensile strength should be converted into a "specific total tensile strength" defined as the sum of the tensile strength measured in the machine and cross machine directions, divided by the basis weight, and corrected in units to a value in meters.

#### **Viscosity**

#### **Overview**

Viscosity is measured at a shear rate of 100 (s<sup>-1</sup>) using a rotational viscometer. The samples are subjected to a linear stress sweep, which applies a range of stresses, each at a constant amplitude.

#### **Apparatus**

Viscometer

Dynamic Stress Rheometer Model SR500 which is available from

15

Sample Plates

25 mm parallel insulated plates are used

Rheometrics Scientific, Inc. of Piscatawy, NJ

Setup

Gap

0.5 mm

Sample Temperature

20°C

20 Sample Volume

at least 0.2455 cm<sup>3</sup>

**Initial Shear Stress** 

10 dynes/cm<sup>2</sup>

**Final Shear Stress** 

1,000 dynes/cm<sup>2</sup>

Stress Increment

25 dynes/cm<sup>2</sup> applied every 20 seconds

#### Method

Place the sample on the sample plate with the gap open. Close the gap and operate the rheometer according to the manufacturer's instructions to measure viscosity as a function of shear stress between the initial shear stress and the final shear stress using the stress increment defined above.

#### Results and Calculation

The resulting graphs plot log shear rate (s<sup>-1</sup>) on the x-axis, log viscosity, Poise (P) on the left y-axis, and stress (dynes/cm<sup>2</sup>) on the right y-axis. Viscosity values are read at a shear rate of 100 (s<sup>-1</sup>). The values for viscosity are converted from P to centipoise (cP) by multiplying by 100.

The disclosures of all patents, patent applications (and any patents which issue thereon, as well as any corresponding published foreign patent applications), and publications mentioned throughout this description are hereby incorporated by reference herein. It is expressly not admitted, however, that any of the documents incorporated by reference herein teach or disclose the present invention.

While particular embodiments of the present invention have been illustrated and described, it would be obvious to those skilled in the art that various other changes and modifications can be made without departing from the spirit and scope of the invention. It is therefore intended to cover in the appended claims all such changes and modifications that are within the scope of this invention.

15

10

What is claimed is:

1. A soft tissue paper product, said soft tissue paper product characterized in that it comprises:

- a) one or more plies of a tissue paper; and
- b) a chemical softening composition deposited on at least one outer surface of said tissue, said chemical softening composition comprising:
  - i) a softening active ingredient, wherein said softening active ingredient comprises a quaternary ammonium compound;
  - ii) an electrolyte, preferably used at a level between 1% and 2% of said softening composition; and
  - iii) a bilayer disrupter; preferably used at a level between 2% and 15% of said softening composition;

and having an extensional viscosity of at least 5 pascal • seconds.

- 2. The tissue paper of Claim 1 wherein said chemical softening composition is deposited as uniform, discrete surface deposits, spaced apart at a frequency between 5 areas per lineal inch and 100 areas per lineal inch.
- 3. The tissue paper of either of Claims 1 or 2 wherein said quaternary ammonium compound has the formula:

$$(R_1)_{4-m} - N^+ - [(CH_2)_n - Y - R_3]_m X^-$$

wherein

Y is -O-(O)C-, or -C(O)-O-, or -NH-C(O)-, or -C(O)-NH-, preferably -O-(O)C- or -C(O)-O-;

m is 1 to 3, preferably 2;

n is 0 to 4, preferably 2;

each R<sub>1</sub> is a C<sub>1</sub>-C<sub>6</sub> alkyl or alkenyl group, hydroxyalkyl group, hydrocarbyl or substituted hydrocarbyl group, alkoxylated group, benzyl group, or mixtures thereof, preferably methyl;

each  $R_3$  is a  $C_{13}$ - $C_{21}$  alkyl or alkenyl group, hydroxyalkyl group, hydroxarbyl or substituted hydroxarbyl group, alkoxylated group, benzyl group, or mixtures thereof, preferably  $C_{15}$ - $C_{17}$  alkyl or alkenyl; and

X is any softener-compatible anion, preferably selected from the group consisting of chloride and methyl sulfate.

4. The tissue paper of any of Claims 1 to 3 wherein said chemical softening composition further comprises a high polymer, preferably selected from the group consisting of polyacrylamide and its derivatives, acrylic polymers and copolymers, water soluble vinyl polymers, polyvinylpyrrolidone, polyethyleneimine, polyalkylene oxides, and mixtures thereof; and preferably having a weight-average molecular weight of at least 500,000.

- 5. The tissue paper of any of Claims 1 to 4 wherein said electrolyte comprises a salt having an anion selected from the group consisting of halides and simple organic acids and having a cation selected from the group consisting of the organic acid salts of sodium, calcium, and magnesium, preferably a sodium formate.
- 6. The tissue paper of any of Claims 1 to 5 wherein said bilayer disrupter is selected from the group consisting of:
  - a) nonionic surfactants derived from saturated and/or unsaturated primary and/or secondary, amine, amide, amine-oxide fatty alcohol, fatty acid, alkyl phenol, and/or alkyl aryl carboxylic acid compounds having from 6 to 22 carbon atoms in a hydrophobic chain, wherein at least one active hydrogen of said compounds is ethoxylated with  $\leq$  50 ethylene oxide moieties to provide an HLB of from 6 to 20;
  - b) nonionic surfactants with bulky head groups selected from the group consisting of:
    - i) surfactants having the formulas:

wherein Y" = N or O; and each R<sup>5</sup> is selected independently from the following: -H, -OH, -(CH<sub>2</sub>)xCH<sub>3</sub>, -O(OR<sup>2</sup>)<sub>z</sub>-H, -OR<sup>1</sup>, - OC(O)R<sup>1</sup>, and -CH(CH<sub>2</sub>-(OR<sup>2</sup>)<sub>z</sub>-H)-CH<sub>2</sub>-(OR<sup>2</sup>)<sub>z</sub>-C(O) R<sup>1</sup>, x and R<sup>1</sup> are as defined above and  $5 \le z$ , z', and z''  $\le 20$ ; and

ii) polyhydroxy fatty acid amide surfactants of the formula:

$$R^2 - C(0) - N(R^1) - Z$$

wherein: each  $R^1$  is H,  $C_1$ - $C_4$  hydrocarbyl,  $C_1$ - $C_4$  alkoxyalkyl, or hydroxyalkyl;  $R^2$  is a  $C_5$ - $C_{21}$  hydrocarbyl moiety; and each Z is a polyhydroxyhydrocarbyl moiety having a linear hydrocarbyl chain with at least 3 hydroxyls directly connected to the chain, or an ethoxylated derivative thereof; and

c) cationic surfactants having the formula:

$${R^{1}_{m} - Y - [(R^{2}-O)_{z} - H]_{p}}^{+} X^{-}$$

wherein  $R^1$  is selected from the group consisting of saturated or unsaturated, primary, secondary or branched chain alkyl or alkyl-aryl hydrocarbons; said hydrocarbon chain having from 6 to 22 carbon atoms; each  $R^2$  is selected from the following groups or combinations of the following groups:  $-(CH_2)_n$ - and/or  $-[CH(CH_3)CH_2]$ -; Y is selected from the following groups:  $= N^+$ - $(A)_q$ ;  $-(CH_2)_n$ - $N^+$ - $(A)_q$ ; -B- $(CH_2)_n$ - $N^+$ - $(A)_2$ ; -(phenyl)- $N^+$ - $(A)_q$ ; -(B-phenyl)- $N^+$ - $(A)_q$ ; with n being from 1 to 4, wherein each A is independently selected from the following groups:  $-(CH_2)_n$ - $-(CH_2)_n$ -

- 7. A composition for softening an absorbent tissue, said composition comprising:
  - a) an effective amount of a softening active ingredient, preferably comprising at least
     35% of said composition;
  - b) a vehicle wherein said softening active ingredient is dispersed;
  - c) an electrolyte dissolved in said vehicle; and
  - d) a bilayer disrupter,

wherein said electrolyte and said bilayer disrupter cooperate to cause the viscosity of said composition to be less than the viscosity of a bicomponent dispersion of said softening active ingredient in said vehicle, and wherein said composition has an extensional viscosity of at least 5 pascal • seconds.

8. The composition of Claim 7 wherein said softening active ingredient comprises a quaternary ammonium compound, preferably wherein said quaternary ammonium compound has the formula:

$$(R_1)_{4-m} - N^+ - [(CH_2)_n - Y - R_3]_m X^-$$

wherein Y is -O-(O)C-, or -C(O)-O-, or -NH-C(O)-, or -C(O)-NH-, preferably -O-(O)C- or -C(O)-O-;

m is 1 to 3, preferably 2;

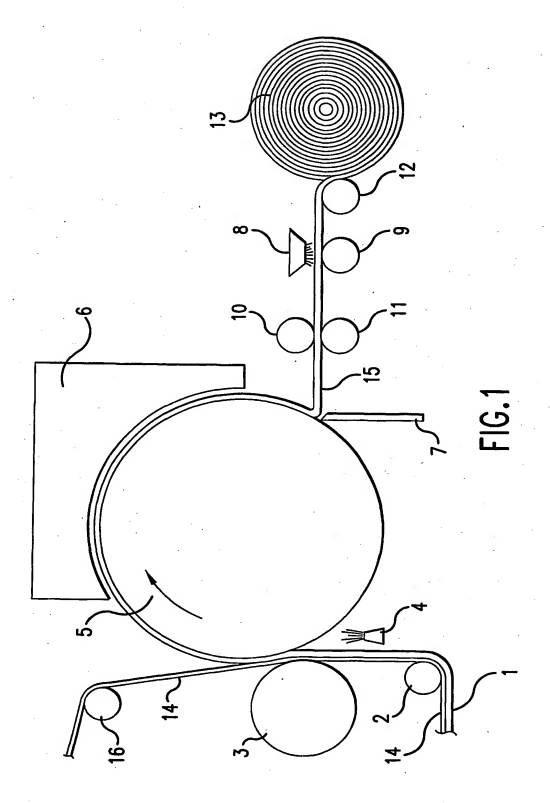
n is 0 to 4, preferably 2;

each  $R_1$  is a  $C_1$ - $C_6$  alkyl or alkenyl group, hydroxyalkyl group, hydroxarbyl or substituted hydroxarbyl group, alkoxylated group, benzyl group, or mixtures thereof, preferably methyl;

each  $R_3$  is a  $C_{13}$ - $C_{21}$  alkyl or alkenyl group, hydroxyalkyl group, hydroxarbyl or substituted hydroxarbyl group, alkoxylated group, benzyl group, or mixtures thereof, preferably  $C_{15}$ - $C_{17}$  alkyl or alkenyl; and

X is any softener-compatible anion, preferably selected from the group consisting of chloride and methyl sulfate.

- 9. A composition according to either of Claims 7 or 8, said composition further comprising a high polymer, preferably comprising between 0.01% and 5% of said composition, and preferably having a weight-average molecular weight of at least 500,000.
- 10. A composition according to Claim 9 wherein said high polymer is selected from the group consisting of polyacrylamide and its derivatives, acrylic polymers and copolymers, water soluble vinyl polymers, polyvinylpyrrolidone, polyethyleneimine, polyalkylene oxides, and mixtures thereof.



In onal Application No Full US 01/48598

A. CLASSIFICATION OF SUBJECT MATTER IPC 7 D21H17/45 D21H17/71

According to International Patent Classification (IPC) or to both national classification and IPC

#### B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)  $IPC \ 7 \quad D21H$ 

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

PAJ, EPO-Internal, WPI Data

	C. DOCUMENTS CONSIDERED TO BE RELEVANT					
Category °	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.				
Х	WO 00 22231 A (PROCTER & GAMBLE) 20 April 2000 (2000-04-20) claims	1-10				
X	WO 00 22233 A (PROCTER & GAMBLE) 20 April 2000 (2000-04-20) claims	1-10				
<b>A</b>	US 5 573 637 A (AMPULSKI ROBERT S ET AL) 12 November 1996 (1996-11-12) examples	1-10				
Α	EP 0 896 045 A (AKZO NOBEL NV ;EKA CHEMICALS AB (SE)) 10 February 1999 (1999-02-10) the whole document	1-10				
	-/	, , , , , , , , , , , , , , , , , , ,				

X Further documents are listed in the continuation of box C.	Patent family members are listed in annex.		
Special categories of cited documents:  'A' document defining the general state of the art which is not considered to be of particular relevance  'E' earlier document but published on or after the International filing date  'L' document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)  'O' document referring to an oral disclosure, use, exhibition or other means  'P' document published prior to the international filing date but later than the priority date claimed	<ul> <li>"T" later document published after the international filing date or priority date and not in conflict with the application but clted to understand the principle or theory underlying the invention</li> <li>"X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone</li> <li>"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.</li> <li>"&amp;" document member of the same patent family</li> </ul>		
Date of the actual completion of the international search	Date of mailing of the international search report		
10 April 2002	19/04/2002		
Name and mailing address of the ISA  European Patent Office, P.B. 5818 Patentiaan 2	Authorized officer		
NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016	Naeslund, P		

Form PCT/ISA/210 (second sheet) (July 1992)

in onal Application No

C (Continue	POOLINEATE CONCERNO	101/US 01/	48598			
Category °	ation) DOCUMENTS CONSIDERED TO BE RELEVANT  Citation of document, with indication, where appropriate, of the relevant passages	<del></del>				
	Citation of document, with indication, where appropriate, of the relevant passages  Relevant to citation of document, with indication, where appropriate, of the relevant passages					
A	US 5 753 079 A (KEYS ROBERT O ET AL) 19 May 1998 (1998-05-19) the whole document		1-10			
<b>A</b>	US 5 814 188 A (FICKE JONATHAN ANDREW ET AL) 29 September 1998 (1998-09-29) the whole document		1-10			
			. *			
			-			

ormation on patent family members

Int onal Application No Pci/US 01/48598

•				FC1/03 (	11/ 46596
Patent document cited in search report	·	Publication date		Patent family member(s)	Publication date
WO 0022231	A	20-04-2000	AU	6424099 A	01-05-2000
WO 0022231	Λ.	. 20 04 2000	AU	6424199 A	01-05-2000
•			BR	9914577 A	03-07-2001
			BR	9914578 A	03-07-2001
				1323367 T	21-11-2001
•		·	CN		21-11-2001
	•		CN	1323368 T	
			CZ	20011348 A3	12-12-2001
			EP	1131486 A1	12-09-2001
			EP	1121488 A1	08-08-2001
			NO	20011806 A	15-06-2001
•			WO	0022233 A1	20-04-2000
			WO	0022231 A1	20-04-2000
				6424000 A	01-05-2000
WO 0022233	Α	20-04-2000	AU AU	6424099 A 6424199 A	01-05-2000
			BR	9914577 A	03-07-2001
	•				03-07-2001
			BR	9914578 A	
			CN	1323367 T	21-11-2001
			CN	1323368 T	21-11-2001
			CZ	20011348 A3	12-12-2001
			EP	1131486 A1	12-09-2001
			ĒΡ	1121488 A1	08-08-2001
*			NO.	20011806 A	15-06-2001
,			WO	0022233 A1	20-04-2000
		· .	WO	0022231 A1	20-04-2000
					12 02 1000
US 5573637	Α	12-11-1996	AU	686919 B2	12-02-1998
			AU	4246796 A	10-07-1996
			BR	9510280 A	06-01-1998
•			CA	2208067 A1	27-06-1996
			CN	1175295 A ,B	04-03-1998
			CZ	9701878 A3	12-11-1997
			EG	20888 A	31-05-2000
			EP	0799350 A1	08-10-1997
			FI	972619 A	18-06-1997
	•	•	หน้	78000 A2	28-04-1999
			JP	3184536 B2	09-07-2001
				10512928 T	08-12-1998
			JP		
			NO	972797 A	19-08-1997
			TR	960569 A2	21-07-1996
			MO	9619616 A1	27-06-1996
			ZA	9510497 A	19-06-1996
EP 0896045	Α	10-02-1999	EP	0896045 A1	10-02-1999
EF 0090043	^	10 02 1939	CA	2244483 A1	06-02-1999
			EP	0896046 A1	10-02-1999
			US	6228223 B1	08-05-2001
				OLLOLLS DI	
US 5753079	Α	19-05-1998	บร	5686023 A	11-11-1997
•			US	5674832 A	07-10-1997
			ΑU	5671996 A	18-11-1996
			CA	2218725 A1	31-10-1996
	•	•	EP	0822859 A1	11-02-1998
			JP	11505555 T	21-05-1999
			WO	9633800 A1	31-10-1996
			WO		
			HC	61621E2 N	
	•		บร	6162453 A	19-12-2000
·	•		US US US	6162453 A 5977189 A 5958863 A	02-11-1999 28-09-1999

formation on patent family members

In all Application No Full US 01/48598

Patent document cited in search report  US 5814188  A 29-09-1998  AU 730321 B2 01-03-2001 AU 5527698 A 31-07-1998 BR 9713643 A 11-04-2000 CN 1245545 A 23-02-2000 EP 0951602 A1 27-10-1999 HU 9904111 A2 28-03-2000 JP 2001507761 T 12-06-2001 NO 993208 A 30-08-1999 TR 9901512 T2 22-11-1999 WO 9829605 A1 09-07-1998 ZA 9711612 A 25-06-1998						
AU 5527698 A 31-07-1998 BR 9713643 A 11-04-2000 CN 1245545 A 23-02-2000 EP 0951602 A1 27-10-1999 HU 9904111 A2 28-03-2000 JP 2001507761 T 12-06-2001 NO 993208 A 30-08-1999 TR 9901512 T2 22-11-1999 WO 9829605 A1 09-07-1998						
	US 5814188	Α	29-09-1998	AU BR CN EP HU JP NO TR WO	5527698 A 9713643 A 1245545 A 0951602 A1 9904111 A2 2001507761 T 993208 A 9901512 T2 9829605 A1	31-07-1998 11-04-2000 23-02-2000 27-10-1999 28-03-2000 12-06-2001 30-08-1999 22-11-1999 09-07-1998

Form PCT/ISA/210 (patent family annex) (July 1992)